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# Method of Inhibiting Protein Tyrosine Phosphatase 1B and/or T-cell Protein Tyrosine Phosphatase and/or other

This application claims the benefit of U.S. Prov. Appl. 60/156,64, sied 09/29/99.

## 5 Field of the Invention

This invention relates to a method of inhibiting Protein Tyrosine Phosphatase 1B (PTP1B) and/or T-cell Protein Tyrosine Phosphatase (TC-PTP) and/or Protein Tyrosine Phosphatases (PTPases) having an aspartic acid (Asp) in position 48 (PTP1B numbering, Chernoff et al., Proc. Natl. Acad. Sci. USA 87: 2735-2789 (1989)) by exposing such an enzyme to inhibitor compounds, i.e., to compounds possessing certain structural, physico-chemical and spatial characteristics that allow them to interact with specific amino acid residues of the active site (and the vicinity of the active site) of PTP1B and/or TC-PTP and more generally Protein Tyrosine Phosphatases (PTPases) having an aspartic acid (Asp) in position 48. The resulting inhibition of the PTPase enzymatic activity makes these compounds useful for elucidating the function of PTP's e.g., by inhibiting a PTP and observing up-or down-regulation of other proteins. Additionally, such inihibitors serve as early development candidates, development candidates, or prototype drugs-for-treatment of or paliation of diseases and dysfunctions such as diabetes type I and II and obesity, cancer, immune disorders (including-allergy-and-abnormal autoimmunity), and conditions involving disturbances in platelet aggregation as well as infectious diseases. This invention also relates to (I) the design and selection of inhibitors which bind to the active site of PTP1B and/or TC-PTP and/or PTPases having an aspartic acid (Asp) in position 48 (II) the synthesis of said inhibitors, methods for their preparation and (III) to compositions comprising the inhibitor compounds.

# 30 Background of the Invention

Protein phosphorylation is now well recognized as an important mechanism utilized by cells to transduce and regulate signals during different stages of cellular function (Hunter, *Phil. Trans. R. Soc. Lond.* B

353: 583-605 (1998); Chan et al., Annu. Rev. Immunol. 12: 555-592 (1994); Zhang, Curr. Top. Cell. Reg. 35: 21-68 (1997); Matozaki and Kasuga, Cell. Signal. 8: 113-19 (1996); Fischer et al, Science 253:401-6 (1991); Flint et al., EMBO J. 12:1937-46 (1993)). The level of tyrosine phosphorylation is balanced by the opposing action of protein tyrosine kinases and protein tyrosine phosphatases. There are at least two major classes of phosphatases: (1) those that dephosphorylate proteins (or peptides) that contain a phosphate group(s) on a serine or threonine moiety (termed Ser/Thr phosphatases) and (2) those that remove a phosphate group(s) from the amino acid tyrosine (termed protein tyrosine phosphatases or PTPases or PTPs).

The PTPases are a family of enzymes that can be classified into two groups: a) intracellular or nontransmembrane PTPases and b) receptor-type or transmembrane PTPases. In addition, dual-specificity phosphatases and low molecular weight phosphatases are able to dephosphorylate phospho tyrosyl proteins. See, e.g., WO 97/ 39746; WO 97/ 40017; WO 99/ 15529; WO 97/08934; WO 98/ 27065; WO 99/46236; WO 99/46244; WO 99/46267; WO 99/46268 and WO 99/46237.

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Intracellular PTPases: Most known intracellular type PTPases contain a single conserved catalytic phosphatase domain consisting of 220-240 amino acid residues. The regions outside the PTPase domains are believed to play important roles in localizing the intracellular PTPases subcellularly (Mauro, L.J. and Dixon, J.E. TIBS 19: 151-155 (1994)). The first intracellular PTPase to be purified and characterized was PTP1B, which was isolated from human placenta (Tonks et al., J. Biol. Chem. 263: 6722-6730 (1988)). Shortly after, PTP1B was expressed recombinantly (Charbonneau et al., Proc. Natl. Acad. Sci. USA 86: 5252-5256 (1989); Chernoff et al., Proc. Natl. Acad. Sci. USA 87: 2735-2789 (1989)). Other examples of intracellular PTPases include (1) T-cell PTPasel TC-PTP (Cool et al. Proc. Natl. Acad. Sci. USA 86: 5257-5261 (1989)), (2) rat brain PTPase (Guan et al., Proc. Natl. Acad. Sci. USA 87:1501-1502 (1990)), (3) neuronal

several different members of the receptor-type PTPase group. Thus, 5 different PTPases. (3) PTP $\alpha$ , (4) PTP $\beta$ , (5) PTP $\delta$ , (6) PTP $\epsilon$ , and (7) PTPC, were identified in one early study (Krueger et al., EMBO J. 9: 3241-3252 (1990)). Other examples of receptor-type PTPases include (8) PTPy (Barnea et al., Mol. Cell. Biol. 13: 1497-1506 (1995)) which, like PTPt (Krueger and Saito, Proc. Natl. Acad. Sci. USA 89: 7417-7421 (1992)) contains a carbonic anhydrase-like domain in the extracellular region, (9) PTPµ (Gebbink et al., FEBS Letters 290: 123-130 (1991)), (10) PTPκ (Jiang et al., Mol. Cell. Biol. 13: 2942-2951 (1993)). Based on structural differences the receptor-type PTPases may be classified into subtypes (Fischer et al., Science 253: 401-406 (1991)): (I) CD45; (II) LAR, PTPd, (11) PTPσ; (III) PTPβ, (12) SAP-1 (Matozaki et al., J. Biol. Chem. 269: 2075-2081 (1994)), (13) PTP-U2/GLEPP1 (Seimiya et al., Oncogene 10: 1731-1738 (1995); Thomas et al., J. Biol. Chem. 269: 19953-19962 (1994)), and (14) DEP-1; (IV) PTPα, PTPε. All receptor-type PTPases except Type III contain two PTPase domains. Novel PTPases are frequently identified, and it is anticipated that between 100 and more than 500 different species will be found in the human genome.

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PTPases are the biological counterparts to protein tyrosine kinases (PTKs). Therefore, one important function of PTPases is to control, and especially down-regulate, the activity of PTKs. However, a more complex picture of the function of PTPases has emerged. Thus, several studies indicate that some PTPases act as positive mediators of cellular signaling. As an example, the SH2 domain-containing SHP-2 acts as a positive mediator in insulin-stimulated Ras activation (Noguchi et al., Mol. Cell. Biol. 14: 6674-6682 (1994)) and of growth factor-induced mitogenic signal transduction (Xiao et al., J. Biol. Chem. 269: 21244-21248 (1994)), whereas the homologous SHP-1 acts as a negative regulator of growth factor-stimulated proliferation (Bignon and Siminovitch, Clin.Immunol.Immunopathol. 73: 168-179 (1994)).

Another example of PTPases as positive regulators has been provided by studies designed to define the activation of the Src-family of tyrosine

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kinases. In particular, several lines of evidence indicate that CD45 is positively regulating the activation of hematopoietic cells, and that the mechanism of such positive regulation may involve dephosphorylation of the C-terminal tyrosine of Fyn and Lck (Chan *et al.*, *Annu. Rev. Immunol.* 12: 555-592 (1994)).

The association of many PTPases with cell proliferation, tranformation and differentiation has now been established. PTP1B, a phosphatase whose structure was the first PTPase to be elucidated (Barford et al., Science 263:1397-1404 (1994)) has been shown to be involved in insulin-induced oocyte maturation (Flint et al., The EMBO J. 12:1937-46 (1993)) and the overexpression of this enzyme has been -associated breast and ovarian cancers (Weiner, implicated in p185 et al., J. Natl. cancer Inst. 86:372-8 (1994); Weiner et al., Am. J. Obstet. Gynecol. 170:1177-883 (1994)). The association with cancer is on the basis of evidence that overexpression of PTP1B is statistically correlated with increased levels of p185<sup>c-erb B2</sup> in ovarian and breast cancer. The role of PTP1B in the etiology and progression of the disease has not yet been elucidated. Inhibitors of PTP1B therefore would help clarify the role of PTP1B in cancer and in some cases provide therapeutic treatment for certain forms of cancer.

# PTPases: the insulin receptor signaling pathway/diabetes

Insulin is an important regulator of different metabolic processes and plays a key role in the control of blood glucose. Defects related to its synthesis or signaling lead to diabetes mellitus. Binding of insulin to the insulin receptor (IR) causes rapid (auto)phosphorylation of several tyrosine residues in the intracellular part of the β-subunit. Three closely positioned tyrosine residues (the tyrosine-1150 domain) must all be phosphorylated to obtain full activity of the insulin receptor tyrosine kinase (IRTK) which transmits the signal further downstream by tyrosine phosphorylation of other cellular substrates, including insulin receptor substrate-1 (IRS-1) (Wilden *et al.*, *J. Biol. Chem. 267*: 16660-16668 (1992); Myers and White, *Diabetes 42*: 643-650 (1993); Lee and

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Pilch, *Am. J. Physiol.* 266: C319-C334 (1994); White *et al.*, *J. Biol. Chem.* 263: 2969-2980 (1988)). The structural basis for the function of the tyrosine-triplet has been provided by X-ray crystallographic studies of IRTK that showed the tyrosine-1150 domain to be autoinhibitory in its unphosphorylated state (Hubbard *et al.*, *Nature* 372: 746-754 (1994)) and of the activated IRTK (Hubbard, *EMBO J.* 16: 5572-5581 (1997)).

Several studies clearly indicate that the activity of the autophosphorylated IRTK can be reversed by dephosphorylation in vitro (reviewed in Goldstein, Receptor 3: 1-15 (1993); Mooney and Anderson, J. Biol. Chem. 264: 6850-6857 (1989)), with the triphosphorylated tyrosine-1150 domain being the most sensitive target for protein-tyrosine phosphatases (PTPases) as compared to the diand mono- phosphorylated forms (King et al., Biochem. J. 275: 413-418) (1991)). This tyrosine-triplet functions as a control switch of IRTK activity and IRTK appears to be tightly regulated by PTP-mediated dephosphorylation in vivo (Khan et al., J. Biol. Chem. 264: 12931-12940 (1989); Faure et al., J. Biol. Chem. 267: 11215-11221 (1992); Rothenberg et al., J. Biol. Chem. 266: 8302-8311 (1991)). The intimate coupling of PTPases to the insulin signaling pathway is further evidenced by the finding that insulin differentially regulates PTPase activity in rat hepatoma cells (Meyerovitch et al., Biochemistry 31: 10338-10344 (1992)) and in livers from alloxan diabetic rats (Boylan et al., J. Clin. Invest. 90: 174-179 (1992)).

Until recently, relatively little was known about the identity of the PTPases involved in IRTK regulation. However, the existence of PTPases with activity towards the insulin receptor can be demonstrated as indicated above. Further, when the strong PTPase-inhibitor pervanadate is added to whole cells an almost full insulin response can be obtained in adipocytes (Fantus et al., Biochemistry 28: 8864-8871 (1989); Eriksson et al., Diabetologia 39: 235-242 (1995)) and skeletal muscle (Leighton et al., Biochem. J. 276: 289-292 (1991)). In addition, other studies show that a new class of peroxovanadium compounds act as potent hypoglycemic compounds in vivo (Posner et al., supra). Two

of these compounds were demonstrated to be more potent inhibitors of dephosphorylation of the insulin receptor than of the EGF-receptor, thus indicating that even such relatively unselective inhibitors may show some specificity in regulating different signal transduction pathways.

It was recently found that mice lacking the protein tyrosine phosphatase-1B gene (PTP1B) (Elchebly *et al.*, *Science 283:* 1544-1548 (1999)) yielded healthy mice thatshowed increased insulin sensitivity and were resistant to diet-induced obesity. These results were confirmed by Kaman at al *Mol. Cell Biol.* 20:5479-5489 (2000). The enhanced insulin sensitivity of the PTP<sup>-/-</sup> mice was also evident in glucose and insulin tolerance tests.

The PTP-1B knock-out mouse showed many characteristics which would be highly desirable results for an anti-diabetes treatment. Most importantly, the knock-out mice grew normally and were fertile and have exhibited no increased incidence of cancer. Blood glucose and insulin levels were lowered, and insulin sensitivity increased. Moreover, the insulin-stimulated tyrosine phosphorylation levels of IR and IRS-1 were found to be increased/prolonged in muscle and liver – but not in fat tissue. Thus, the main target tissues for this type of approach would appear to be insulin action in liver and muscle.

Several other "diabetic" parameters were also improved, including plasma triglycerides which were decreased in the knock-out mice. The knock-animals also exhibited a resistance to weight gain when placed on a high-fat diet. This is in contrast to the action of the PPARγ agonist class of insulin sensitizers, which rather induce weight gain (Murphy & Nolan, *Exp. Opin. Invest. Drugs* 9:1347-1361, 2000), and would suggest that inhibition of PTP-1B could be a particularly attractive option for treatment of obese Type II diabetics.

This is also supported by the fact that the heterozygous mice from this study showed many of these desirable features. The reduction in weight gain of the knock-out animals on the high fat diet was found to be due to a decreased fat cell mass, although differences were observed with respect to fat cell number. Leptin levels were also lower in the knock-out

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mice, presumably as a reflection of the decreased fat mass. Significantly, the Klaman et al group also found that the knock-out animals had an increased energy expenditure of around 20% and an increased respiratory quotient compared to the wild-type; again, heterozygote animals displayed an intermediate level of energy expenditure. Therefore, inhibition of this enzyme may be an effective anti-diabetic and perhaps also anti-obesity therapy.

It should also be noted that in the PTP-1B knock-out mice the basal tyrosine phosphorylation level of the insulin receptor tyrosine kinase does not appear to be increased, which is in contrast to the situation after insulin treatment where there is an increased or prolonged phosphorylation. This might indicate that other PTPs are controlling the basic phosphorylation state of the insulin receptor in the knock-out mice – and is expected to do so in man.

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Also other PTPases have been implicated as regulators of the insulin signaling pathway. Thus, it was found that the ubiquitously expressed SH2 domain containing PTPase, PTP1D/SHP-2 (Vogel et al., 1993, supra), associates with and dephosphorylates IRS-1, but apparently not the IR itself (Kuhné et al., J. Biol. Chem. 268: 11479-11481 (1993); (Kuhné et al., J. Biol. Chem. 269: 15833-15837 (1994)).

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Other studies suggest that receptor-type or membrane-associated PTPases are involved in IRTK regulation (Faure *et al., J. Biol. Chem.* 267: 11215-11221 (1992), (Häring *et al., Biochemistry* 23: 3298-3306 (1984); Sale, *Adv. Prot. Phosphatases* 6: 159-186 (1991)).

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While previous reports indicate a role of PTP $_{\alpha}$  in signal transduction through src activation (Zheng *et al.*, *Nature 359*: 336-339 (1992); den Hertog *et al.*, *EMBO J. 12*: 3789-3798 (1993)) and interaction with GRB-2 (den Hertog *et al.*, *EMBO J. 13*: 3020-3032 (1994); Su *et al.*, *J. Biol. Chem. 269*: 18731-18734 (1994)), Møller, Lammers and coworkers provided results that suggest a function for this phosphatase and its close relative PTP $_{\epsilon}$  as negative regulators of

the insulin receptor signal (Møller et al., 1995 supra;

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Lammers, et al., FEBS Lett. 404:37-40 (1997). These studies also indicated that receptor-like PTPases may play a significant role in regulating the IRTK, including through direct influence on the insulin receptor itself.

Other studies have shown that PTP1B and TC-PTP are likely to be involved in the regulation of several other cellular processes in addition to the described regulatory roles in insulin signaling. Therefore, PTP1B and/or TC-PTP as well as other PTPases showing key structural features with PTP1B and TC-PTP are likely to be important therapeutic targets in a variety of human and animal diseases. The compounds of the present invention are useful for modulating or inhibiting PTP1B and/or TC-PTP and/or other PTPases showing key structural features with said PTPases and thus elucidating their function and for treating disease states in which said modulation or inhibition is indicated.

Further, PTPases influence the following hormones or diseases or disease states: somatostatin, the immune system/autoimmunity, cell-cell interactions/cancer, platelet aggregation, osteoporosis, and microorganisms, as disclosed in PCT Publication WO 99/15529.

## 20 PTPases: the immune system/autoimmunity

Several studies suggest that the receptor-type PTPase CD45 plays a critical role not only for initiation of T cell activation, but also for maintaining the T cell receptor-mediated signaling cascade. These studies are reviewed in: (Weiss A., *Ann. Rev. Genet. 25*: 487-510 (1991); Chan *et al.*, *Annu. Rev. Immunol. 12*: 555-592 (1994); Trowbridge and Thomas, *Annu. Rev. Immunol. 12*: 85-116 (1994)).

CD45 is one of the most abundant of the cell surface glycoproteins and is expressed exclusively on hemopoetic cells. In T cells, it has been shown that CD45 is one of the critical components of the signal transduction machinery of lymphocytes. In particular, there is evidence that CD45 phosphatase plays a pivotal role in antigen-stimulated proliferation of T lymphocytes after an antigen-has-bound to the T cell receptor (Trowbridge, *Ann. Rev. Immunol*, 12: 85-116 (1994)). Several studies indicate that the PTPase activity of CD45 plays a role in the

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activation of Lck, a lymphocyte-specific member of the Src family proteintyrosine kinase (Mustelin etal., Proc. Natl. Acad. Sci. USA 86: 6302-6306 (1989); Ostergaard et al., Proc. Natl. Acad. Sci. USA 86: 8959-8963 (1989)). Studies using transgenic mice with a mutation for the CD45-5 exon6 exhibited a lack of mature T cells. These mice did not respond to an antigenic challenge with the typical T cell mediated response (Kishihara et al., Cell 74:143-56 (1993)). Inhibitors of CD45 phosphatase would therefore be very effective therapeutic agents in conditions that are associated with autoimmune diseases, such as rheumatoid arthritis, systemic lupus erythematosus, type I diabetes, and inflammatory bowel disease. Another important function of CD45 phosphatase inhibitors is in effecting immunosuppression, where such a result is indicated, e.g., in transplantation and other conditions in need of immunosuppressive treatment.

CD45 has also been shown to be essential for the antibody mediated degranulation of mast cells (Berger et al., J. Exp. Med. 180:471-6 (1994)). These studies were also done with mice that were CD45deficient. In this case, an IgE-mediated degranulation was demonstrated in wild type but not CD45-deficient T cells from mice. These data suggest that CD45 inhibitors could also play a role in the symptomatic or therapeutic treatment of allergic disorders, such as asthma, allergic rhinitis, food allergies, eczema, urticaria and anaphylaxis. Another PTPase, an inducible lymphoid-specific protein tyrosine phosphatase (HePTP) has also been implicated in the immune response. This phosphatase is expressed in both resting T and B lymphocytes, but not non-hemopoetic cells. Upon stimulation of these cells, mRNA levels from the HePTP gene increase 10-15 fold (Zanke et al., Eur. J. Immunol. 22: 235-239 (1992)).

Likewise, the hematopoietic cell specific SHP-1 acts as a negative regulator and thus appears to play an essential role in immune cell development. In accordance with the above-mentioned important function of CD45, HePTP and SHP-1, selective PTPase inhibitors are early development candidates or prototype drugs both as immunosuppressors and as immunostimulants. Recent studies illustrate the potential of

PTPase inhibitors as immunmodulators by demonstrating the capacity of the vanadium-based relatively nonselective PTPase inhibitor, BMLOV, to induce apparent B cell selective apoptosis compared to T cells (Schieven et al., J. Biol. Chem. 270: 20824-20831 (1995)).

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## PTPases: cell-cell interactions/cancer

Focal adhesion plaques, an in vitro phenomenon in which specific contact points are formed when fibroblasts grow on appropriate substrates, mimic, in certain respects, cells and their natural surroundings. Several focal adhesion proteins are phosphorylated on tyrosine residues when fibroblasts adhere to and spread on extracellular matrix (Gumbiner, Neuron 11: 551-564 (1993)). However, aberrant tyrosine phosphorylation of these proteins can lead to cellular transformation. The intimate association between PTPases and focal adhesions is supported by the finding of several intracellular PTPases with ezrin-like N-terminal domains, e.g. PTPMEG1 (Gu et al., Proc. Natl. Acad. Sci. USA 88: 5867-5871 (1991), PTPH1 (Yang and Tonks, Proc. Natl. Acad. Sci. USA 88: 5949-5953 (1991)) and PTPD1 (Møller et al., Proc. Natl. Acad. Sci. USA 91: 7477-7481 (1994)). The ezrin-like domains show similarity to several proteins that are believed to act as links between the cell membrane and the cytoskeleton. PTPD1 was found to be phosphorylated by and associated with c-src in vitro and is hypothesized to be involved in the regulation of phosphorylation of focal adhesions (Møller et al., supra).

PTPases may oppose the action of tyrosine kinases, including those responsible for phosphorylation of focal adhesion proteins, and may therefore function as natural inhibitors of transformation. TC-PTP, and especially the truncated form of this enzyme (Cool *et al.*, *Proc. Natl. Acad. Sci. USA 87*: 7280-7284 (1990)), can inhibit the transforming activity of v-*erb* and v-*fms* (Lammers *et al.*, *J. Biol. Chem. 268*: 22456-22462 (1993), Zander *et al.*, *Oncogene 8*: 1175-1182 (1993)). Moreover, it was found that transformation by the oncogenic form of the *HER2/neu* gene was suppressed in NIH 3T3 fribroblasts

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overexpressing PTP1B (Brown-Shimer et al., Cancer Res. 52: 478-482 (1992)).

The expression level of PTP1B was found to be increased in a mammary cell line transformed with *neu* (Zhay *et al., Cancer Res. 53:* 2272-2278 (1993)). The intimate relationship between tyrosine kinases and PTPases in the development of cancer is further evidenced by the recent finding that PTPe is highly expressed in murine mammary tumors in transgenic mice over-expressing c-*neu* and v-Ha-*ras*, but not c-*myc* or *int-2* (Elson and Leder, *J. Biol. Chem. 270:* 26116-26122 (1995)). Further, the human gene encoding PTPγ was mapped to 3p21, a chromosomal region which is frequently deleted in renal and lung carcinomas (LaForgia *et al., Proc. Natl. Acad. Sci. USA 88:* 5036-5040 (1991)).

PTPases appear to be involved in controlling the growth of fibroblasts. In a recent study it was found that Swiss 3T3 cells harvested at high density contain a membrane-associated PTPase whose activity on an average is 8-fold higher than that of cells harvested at low or medium density (Pallen and Tong, *Proc. Natl. Acad. Sci. USA 88*: 6996-7000 (1991)).

Two closely related receptor-type PTPases, PTP $_{\rm K}$  and PTP $_{\rm H}$ , can mediate homophilic cell-cell interaction when expressed in non-adherent insect cells, suggesting that a normal physiological function for these PTPases in cell-to-cell signalling (Gebbink *et al., J. Biol. Chem. 268*: 16101-16104 (1993), Brady-Kalnay *et al., J. Cell Biol. 122*: 961-972 (1993); Sap *et al., Mol. Cell. Biol. 14*: 1-9 (1994)). Interestingly, PTP $_{\rm K}$  and PTP $_{\rm H}$  do not bind to each other (PTP $_{\rm K}$  does self-associate), despite their structural similarity (Zondag *et al., J. Biol. Chem. 270*: 14247-14250 (1995)).

From the studies described above it is apparent that PTPases
play an important role in regulating normal cell growth. Additionally, as pointed out above, PTPases may also function as positive mediators of intracellular signaling and thereby induce or enhance mitogenic responses. Increased activity of certain PTPases might therefore result

in cellular transformation and tumor formation. See, Zheng, supra; Uchida et al., J. Biol. Chem. 269: 12220-12228 (1994 Hunter, Cell 80: 225-236 (1995). Inhibitors of specific PTPases are therefore likely to be of significant therapeutic value in the treatment of certain forms of cancer.

# PTPases: platelet aggregation

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PTPases are centrally involved in platelet aggregation. Thus, agonist-induced platelet activation results in calpain-catalyzed cleavage of PTP1B with a concomitant 2-fold stimulation of PTPase activity (Frangioni *et al.*, *EMBO J. 12*: 4843-4856 (1993)). The cleavage of PTP1B leads to subcellular relocation of the enzyme and correlates with the transition from reversible to irreversible platelet aggregation in platelet-rich plasma. In addition, the SH2 domain containing PTPase, SHP-1, was found to translocate to the cytoskeleton in platelets after thrombin stimulation in an aggregation-dependent manner (Li *et al.*, *FEBS Lett. 343*: 89-93 (1994)).

Although some details in the above two studies have been questioned, there is overall agreement that PTP1B and SHP-1 play significant functional roles in platelet aggregation (Ezumi *et al., J. Biol. Chem. 270:* 11927-11934 (1995)). In accordance with these observations, treatment of platelets with the PTPase inhibitor pervanadate leads to significant increase in tyrosine phosphorylation, secretion and aggregation (Pumiglia *et al., Biochem. J. 286:* 441-449 (1992)).

### PTPases: osteoporosis

The rate of bone formation is determined by the number and the activity of osteoblasts. In turn, these are determined by the rate of proliferation and differentiation of osteoblast progenitor cells, respectively. Histomorphometric studies indicate that the osteoblast number is the primary determinant of the rate of bone formation inhumans (Gruber et al., Mineral Electrolyte Metab. 12: 246-254 (1987), reviewed in Lau et al., Biochem. J. 257: 23-36 (1989)). Acid

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phosphatases/PTPases are implicated in negative regulation of osteoblast proliferation. Thus, fluoride, which has phosphatase inhibitory activity, has been found to increase spinal bone density in osteoporotics by increasing osteoblast proliferation (Lau *et al.*, *supra*).

Consistent with this observation, an osteoblastic acid phosphatase with PTPase activity was found to be highly sensitive to mitogenic concentrations of fluoride (Lau et al., J. Biol. Chem. 260: 4653-4660 (1985), Lau et al., J. Biol. Chem. 262: 1389-1397 (1987), Lau et al., Adv. Protein Phosphatases 4: 165-198 (1987)). The mitogenic action of fluoride and other phosphatase inhibitors (molybdate and vanadate)

may thus be explained by their inhibition of acid phosphatases/PTPases that negatively regulate the cell proliferation of osteoblasts. The complex nature of the involvement of PTPases in bone formation is further suggested by the recent identification of a

novel parathyroid regulated, receptor-like PTPase, OST-PTP, expressed in bone and testis (Mauro et al., J. Biol. Chem. 269: 30659-30667 (1994)). OST-PTP is up-regulated following differentiation and matrix formation of primary osteoblasts and subsequently down-regulated in the osteoblasts which are actively mineralizing bone in culture. In addition, it was recently observed that vanadate, vanadyl and

pervanadate all increased the growth of the osteoblast-like cell line
UMR106. Vanadyl and pervanadate were stronger stimulators of cell
growth than vanadate. Only vanadate was able to regulate the cell
differentiation as measured by cell alkaline phosphatase activity

(Cortizo et al., Mol. Cell. Biochem. 145: 97-102 (1995)). More important, several studies have shown that biphosphonates, such as alendronate and tiludronate, inhibit PTPase activity in osteoclasts and that the inhibition of PTPase activity correlated with the inhibition of *in vitro* osteoclast formation and bone resorption (Scmidt, et al., *Proc. Natl Acad. Sci. U.S.A.* 93: 3068-3073,1996; Murakami et al., *Bone 20*:399-

Acad. Sci. U.S.A. 93: 3068-3073,1996; Murakami et al., Bone 20:399-404, 1997; Opas et al., Biochem. Pharmacol. 54: 721-727, 1997; Skorey et al., J. Biol. Chem. 272: 22472-22480, 1997. Thus, other PTPase inhibitors are potentially effective in countering osteoclast activity, and thus treating osteoporosis.

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# PTPases: microorganisms

Dixon and coworkers have called attention to the fact that PTPases may be a key element in the pathogenic properties of Yersinia (reviewed in Clemens et al. Molecular Microbiology 5: 2617-2620 (1991)). This finding was rather surprising since tyrosine phosphate is thought to be absent in bacteria. The genus Yersinia comprises 3 species: Y. pestis (responsible for the bubonic plague), Y. pseudoturberculosis and Y. enterocolitica (causing enteritis and mesenteric lymphadenitis). A dual-specificity phosphatase, VH1, has 10 been identified in Vaccinia virus (Guan et al., Nature 350: 359-263 (1991)). These observations indicate that PTPases may play critical roles in microbial and parasitic infections, and they further point to PTPase inhibitors as a novel, putative treatment principle of infectious diseases. Availibility of PTPase inhibitors would help shed light in all 15 the foregoing specualations about PTPase function because they would enable assaying techniques which would answer some of these questions as will be illustrated below.

# 20 Summary of Background

It has been found that PTPases play a major role in the above modulation and regulation of fundamental cellular signaling mechanisms involved in metabolism, growth, proliferation and differentiation (Fisher *et al*, *Science* 253:401-6 (1991); Tonks and Neel, *Cell* 87: 365-368 (1966)" Neel and Tonks, *Current Opinion in Cell Biology* 9: 193-204 (1997); Hunter, *Phil. Trans. R. Soc. Lond. B* 353: 583-605 (1998); Hunter, *Cell* 100: 113-120 (2000); Zhang, *Critical Reviews in Biochemistry and Molecular Biology* 33:1-52 (1988)). Reports from many laboratories have shown that PTPases can act both as positive and negative regulators of signal transduction processes. PTPases have been implicated in a variety of human diseases, including diabetes, obesity, autoimmune diseases, acute and chronic inflammation, osteoporosis, proliferative-disorders including various forms of cancer, growth disorders, and defective platelet

aggregation (WO97/39748, WO97/40017, WO99/1529, WO97/08934, WO98/27065, WO99/46236, WO99/46244, WO99/46267, WO99/46268, WO99/46237). Accordingly there is increasing evidence which suggests that inhibition of these PTPases would help treat or manage these diseases (Hunter, *vide supra*; Neel and Tonks, *vide supra*: Frangione et al., *EMBO J.* 12:4843-4856 (1993); Zhang, *Curr. Top. Cell. Reg.* 35. 21-68 (1997): Zhang, *vide supra*; Evans and Jalian, *Exp. Opinion. Invest. Drugs* 8: 139-160 (1999); Burke and Zhang, *Bioploymers (Peptide Science)* 47: 225-241 (1998): Elchebly *et al.*; *Science* 283: 1544-1548 (1999); Wrobel et al.; *J. Med. Chem.* 42: 3199-3202 (1999)). In addition, certain infectious diseases may also be treated or managed by administration PTPase inhibitors (Clemens *et al., Molecular Microbiology* 5: 2617-2620 (1991)).

Both selective PTPase inhibitors and inhibitors that bind to several PTPases (non-selective inhibitors) can be used therapeutically to partially or completely restore PTPase-mediated perturbed signal transduction processes and thus for management, treatment, palliation or prevention of the above diseases.

## 20 **Description of Drawings**

Figure 1. Active site of Protein Tyrosine Phosphatase 1B complexed with with 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid.

- Figure 2. Active site of Protein Tyrosine Phosphatase 1B complexed with 7-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (Example 26).
- Figure 3. Active site of Protein Tyrosine Phosphatase 1B complexed with 5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (Example 4).

Figure 4. Active site of Protein Tyrosine Phosphatase 1B complexed with 2-(oxalyl-amino)-7-(1,1,3-trioxo-1H-benzo[d]isothiazol-3-yloxomethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (Example 54). Selected water molecules are shown.

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## **Description of the Invention**

The present invention provides a method for inhibiting protein tyrosine phosphatase 1B (PTP1B) and/or T-cell protein tyrosine phosphatase (T-cell PTP/TC-PTP) and/or protein tyrosine phosphatases (PTPases) having an aspartic acid (Asp) in position 48 by exposing said PTPase to a compound having physico-chemical and spatial structural characteristics that interfere with the active site and/or vicinity of the active site of said PTPase thereby inhibiting its enzymatic activity. Specifically, the present inhibitors of PTP1B and/or TC-PTP and/or PTPases having an aspartic acid (Asp) in position 48 interact with two or more residues of the following: arginine 221, glycine 220, lysine 120, tyrosine 46, and phenylalanine/histidine 182 and one or more of the following (residue numbering correspondingto PTP1B will be used through out (Chernoff et al., Proc. Natl. Acad. Sci. USA 87: 2735-2789 (1989)):

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- 1. Isoleucine 219 backbone amide nitrogen;
- 2. Glycine 218 backbone amide nitrogen;
- 3. Alanine 217 backbone amide nitrogen;
- 4. Serine 216 backbone amide nitrogen;
- 5. Cysteine 215 backbone amide nitrogen;
- 25 6. The side chain carboxylic acid group of aspartic acid 181;
  - 7. The side chain carboxylic acid group of aspartic acid 48;
  - 8. The side chain guanidinium group of arginine 47;
  - 9. Arginine 47 backbone amide nitrogen;
  - 10. Aspartic acid 48 backbone amide nitrogen;
  - 11. The side chain hydroxy group of tyrosine 46;
    - 12. The side chain amino group of lysine 41;
    - 13. The methylene-side chain atoms of lysine 41;
    - 14. The backbone amide carbonyl of asparagine 44;
    - 15. The methylene side chain atoms of arginine 45;

- 16. The backbone amide carbonyl of arginine 45;
- 17. The methylene side chain atoms of arginine 47;
- 18. The methylene side chain atom of aspartic acid 48;
- 19. The backbone amide carbonyl of aspartic acid 48;
- 5 20. The methylene side chain atoms of leucine 88;
  - 21. The side chain hydroxy group of serine 118;
  - 22. The backbone amide carbonyl of leucine 119;
  - 23. The side chain amide nitrogen of glutamine 262;
  - 24. The side chain atoms of methionine 258;
- 10 25. The aromatic group of phenylalanine 52;
  - 26. The backbone amide nitrogen of glycine 259;
  - 27. The alpha-methylene atom of glycine 259;
  - 28. The guanidinium group of arginine 254;
  - 29. The methylene side chain atoms of arginine 254;
- 15 30. The methylene side chain atoms of arginine 24;
  - 31. The guanidinium group of arginine 24; or
  - 32. Any conserved water molecule in the vicinity of the active site.

Preferably, the present inhibitors of PTP1B and/or TC-PTP and/or PTPases having an aspartic acid (Asp) in position 48 interact with any three or more of the above identified regions of the active site and its vicinity.

In one preference, the inhibitors of PTP1B and/or TC-PTP and/or

PTPases having an aspartic acid (Asp) in position 48 interact with arginine

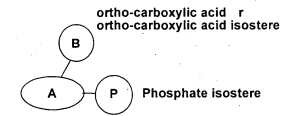
221, glycine 220, lysine 120, tyrosine 46, phenylalanine/histidine 182,

aspartic acid 48 and one or more of the following

- 1. Isoleucine 219 backbone amide nitrogen;
- 2. Glycine 218 backbone amide nitrogen;
- 3. Alanine 217 backbone amide nitrogen;
  - 4. Serine 216 backbone amide nitrogen;
  - The side chain carboxylic acid group of aspartic acid-181;
  - The side chain quanidinium group of arginine 47;
  - Arginine 47 backbone amide nitrogen;

- 8. Aspartic acid 48 backbone amide nitrogen;
- 9. The side chain hydroxy group of tyrosine 46;
- 10. The side chain amino group of lysine 41;
- 11. The methylene side chain atoms of lysine 41;
- 12. The backbone amide carbonyl of asparagine 44;
  - 13. The methylene side chain atoms of arginine 45;
  - 14. The backbone amide carbonyl of arginine 45;
  - 15. The methylene side chain atoms of arginine 47;
  - 16. The methylene side chain atom of aspartic acid 48;
- 10 17. The backbone amide carbonyl of aspartic acid 48;
  - 18. The methylene side chain atoms of leucine 88;
  - 19. The side chain hydroxy group of serine 118;
  - 20. The backbone amide carbonyl of leucine 119;
  - 21. The side chain amide nitrogen of glutamine 262;
- 15 22. The side chain atoms of methionine 258;
  - 23. The aromatic group of phenylalanine 52;
  - 24. The backbone amide nitrogen of glycine 259;
  - 25. The alpha-methylene atom of glycine 259;
  - 26. The guanidinium group of arginine 254;
- 20 27. The methylene side chain atoms of arginine 254;
  - 28. The methylene side chain atoms of arginine 24;
  - 29. The guanidinium group of arginine 24; or
  - 30. Any conserved water molecule in the vicinity of the active site.
- Preferred key structural features of the inhibitors of the present invention include a phosphate isostere (**P**), a carboxylic acid perferably or a carboxylic acid or ortho-carboxylic acid or o-c acid isostere (**B**) and a hydrophobic group (**A**) as shown in Scheme 1.

#### Scheme 1.



Hydrophobic group

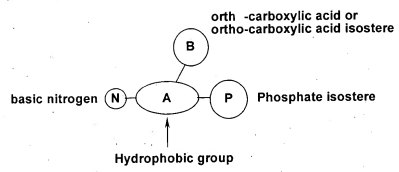
In a preferred embodiment, the key structural features of the inhibitors of the present invention include a phosphate isostere (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B) and a hydrophobic group (A), preferably a phenyl, naphthyl or thiophenyl as shown in Scheme 1.

In another preferred embodiment the key structural features of the inhibitors of the present invention include an oxalylamide (-NHCOCOOH) (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B) and a hydrophobic group (A).

In another preferred embodiment the key structural features of the
inhibitors of the present invention include an oxalylamide (-NHCOCOOH)
(P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B) and a hydrophobic group (A), preferably a phenyl, naphthyl or thiophenyl as shown in Scheme 1.

In another preferred embodiment the key structural features of the inhibitors of the present invention include a phosphate isostere (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), a hydrophobic group (A) and a basic nitrogen (N) as shown in Scheme 2.

## Scheme 2.



- In another preferred embodiment, the key structural features of the inhibitors of the present invention include an oxalylamide (-NHCOCOOH) (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), a hydrophobic group (A) and a basic nitrogen (N) as shown in Scheme 2.
- In another preferred embodiment, the key structural features of the inhibitors of the present invention include an oxalylamide (-NHCOCOOH) (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), a hydrophobic group (A), preferably a phenyl, naphthyl or thiophenyl and a basic nitrogen (N).

In another preferred embodiment, the key structural features of the inhibitors of the present invention include a basic nitrogen which provides selectivity for PTPases containing an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - a phosphate isostere (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), a hydrophobic group (A) as shown in Scheme 3.

## Scheme 3.

In another preferred embodiment, the key structural features of the inhibitors of the present invention include a basic nitrogen which provides selectivity for PTPases containing an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - an oxalylamide (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), a hydrophobic group (A) as shown in Scheme 3.

In another preferred embodiment, the key structural features of the inhibitors of the present invention include a basic nitrogen which provides selectivity for PTPases containing an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - an oxalylamide (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), and a hydrophobic group (A), preferably a phenyl, naphthyl or thiophenyl as shown in Scheme 3.

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## Scheme 4.

In another preferred embodiment, the key structural features of the inhibitors of the present invention include a basic nitrogen which provides selectivity for PTPases containing an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - a phosphate isostere (P), an ortho-carboxylic acid or ortho-carboxylic acid isostere (B), an aromatic group (A), preferably a phenyl or thiophenyl and a hydrophobic group (H) as shown in Scheme 4.

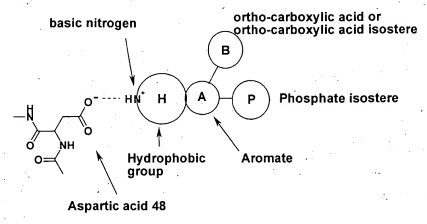
In another preferred embodiment, the key structural features of the inhibitors of the present invention include a basic nitrogen which provides selectivity for PTPases containing an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - an oxalylamide (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), an aromatic group (A), preferably a phenyl or thiophenyl and a hydrophobic group (H) as shown in Scheme 4.

In another preferred embodiment, the key structural features of the inhibitors of the present invention include a phosphate isostere (P), an ortho-carboxylic acid or an ortho-carboxylic acid isostere (B), an aromatic group (A), preferably a phenyl or thiophenyl and a hydrophobic group (H) which include a basic nitrogen which provides selectivity for PTPases that

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contain an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - as shown in Scheme 5.

#### 5 Scheme 5.



In another preferred embodiment, the key structural features of the inhibitors of the present invention include an oxalylamide (P), an orthocarboxylic acid or an orthocarboxylic acid isostere (B), an aromatic group (A), preferably a phenyl or thiophenyl and a hydrophobic group (H) which include a basic which provides selectivity for PTPases that contain an aspartic acid in position 48 - via formation of a salt bridge to said aspartic acid 48 and repulsion to PTPases that contain the corresponding asparagine in position 48 - as shown in Scheme 5.

The key structural features of the inhibitors of the present invention described above are linked to each other via covalent bonds.

The compounds of the present invention possess, but are not limited to, a phosphate isostere in which the centroid of the phosphate isostere is 5.0-5.5 Å from the centroid of a carboxylic acid or carboxylic acid isostere, and 4.5-5.1 Å from the centroid of an aromatic group or a hydrophobic group. In a preferred embodiment, the compounds of the present invention possess, but are not limited to, an oxalylamide in which the centroid of the carboxylic acid moiety of said oxalylamide is 5.0-5.5 Å from the

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centroid of a carboxylic acid or carboxylic acid isostere, and 4.5-5.1 Å from the centroid of an aromatic group or a hydrophobic group. In an other preferred embodiment the compounds of the present invention possess, but are not limited to, a phosphate isostere in which the centroid of the phosphate isostere is 5.0-5.5 Å from the centroid of a carboxylic acid or carboxylic acid isostere, 4.5-5.1 Å from the centroid of an aromatic group or a hydrophobic group and 8.0-14.0 Å from a basic nitrogen. These features must participate in the appropriate interactions (e.g. hydrogen bonds, salt bridges, hydrophobic interactions, cation- $\pi$ 

interactions, or  $\pi$ ,  $\pi$  interactions, or aromatic-aromatic interactions) with the PTP1B and/or TC-PTP and/or other PTPases that are structurally similar to PTP1B active site and vicinity e.g. having an aspartic acid (Asp) in position 48. The centroid of the phosphate isostere should be 3.50-4.20 Å from the centroid of the side chain guanidinium group of arginine 221.

The centroid of the carboxylic acid or carboxylic acid isostere should be 3.4-4.1 Å from the side chain amino group of lysine 120. The basic nitrogen should be 3.4-4.1 Å from the centroid of aspartic acid 48. The aromatic or, more generally, hydrophobic group should be near the following amino acid side chain atoms with appropriate distance ranges between the centroid of the side chain atoms and the centroid of the aromatic - or hydrophobic group given in parentheses: tyrosine 46 (4.4-5.1 Å) and phenylalanine/histidine 182 (4.4-6.5 Å).

The centroid of the oxalylamide carboxylic acid moiety should be 3.50-4.20 Å from the centroid of the side chain guanidinium group of arginine 221. The centroid of the carboxylic acid or carboxylic acid isostere should be 3.4-4.1 Å from the side chain amino group of lysine 120. The basic nitrogen should be 3.4-4.1 Å from the centroid of aspartic acid 48. The aromatic - or hydrophobic group should be near the following amino acid side chain atoms with appropriate distance ranges between the centroid of the side chain atoms and the centroid of the aromatic - or hydrophobic group given in parentheses: tyrosine 46 (4.4-5.1 Å) and phenylalanine/histidine 182 (4.4-6.5 Å).

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In a specific embodiment, the invention is directed to a method of inhibiting at least one intracellular or membrane-associated PTPase that has aspartic acid (Asp) in position 48 using the numbering for PTP1B, the method comprising exposing the PTPase to an inhibitor compound which fits spatially into the active site and the vicinity thereof, said compound comprising the following features and moieties:

- (a) a phosphate isostere which forms a salt bridge to the guanidinium group of arginine 221 and a hydrogen bond with a hydrogen. atom donated by the backbone amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the phosphate isostere group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 backbone amide nitrogen ranges from 3.5-4.2 Å, and (III) said glycine 220 backbone amide nitrogen ranges from 2.7-3.5 Å; or (b) an oxalylamide which forms a salt bridge to the quanidinium group of arginine 221 and forms a hydrogen bond with a hydrogen atom donated by the amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the carboxylic acid group of said oxalylamide group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 amide nitrogen ranges from 3.5-4.2 Å and the distance between the amide carbonyl group of said oxalylamide group and the said glycine 220 amide nitrogen ranges from 2.7-3.5 Å; and
- II. (a) a carboxylic acid group or (b) a carboxylic acid isostere
  group selected from the following 5-membered heterocycles

wherein said acid or said isostere group forms a salt bridge to the side chain amino group of lysine 120 wherein the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said Lysine 120 ranges from 3.4-4.1 Å; and

III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said

hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å;

and at least one of features IV through V:

- IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 4.4-5.1 Å; and
- 10 V. a hydrophobic group that interacts with the imidazole ring of histidine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said histidine 182 ranges from 4.4-6.5 Å; and

one or more of the following features VI-XXXVII:

- VI. an amino group which forms a salt bridge to the site chain carboxylic acid group of aspartic acid 48 such that the distance between the nitrogen atom of said amino group and the centroid of said site chain carboxylic acid group of aspartic acid 48 ranges from 3.4-4.1 Å; and
- VII. two oxygen atoms which form hydrogen bonds via a water molecule to the side chain carboxylic acid group of aspartic acid 48 such that the distance between each of the two oxygen atoms and the centroid of said water molecule ranges from 2.5-3.6 Å and that the distance between said water molecule and the centroid of said side chain carboxylic acid group of aspartic acid 48 ranges from 2.5-3.6 Å and that the distance between said two oxygen atoms ranges from 2.5-3.0 Å; and

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VIII. a hydrophobic group that interacts with the side chain methylene-groups-of-tyrosine-46-such that the distance between the centroid of said hydrophobic group and the centroid of the methylene groups of said tyrosine 46 ranges from 4.4-5.1 Å;

- IX. a hydrophilic group that forms a hydrogen bond or forms a salt bridge with aspartic acid 181 such that the distance between the centroid of said hydrophilic group and the centroid of the carboxylic acid of said aspartic acid 181 ranges from 4.4-5.1 Å;
- X. a hydrophobic group that interacts with tyrosine 46 and the methylene side chain atoms of arginine 47 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 is 4.7-5.2 Å and the centroid of the methylene side chain atoms of said arginine 47 ranges from 4.5-5.5 Å;
- XI. a hydrophilic group that forms a hydrogen bond with the one or more hydrogen atoms donated by the guanidinium group of arginine 47 such that the distance between the centroid of said hydrophilic group and the guanidinium group of said arginine 47 ranges from 2.7-3.5 Å;
- XII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of arginine 47 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said arginine 47 is 2 ranges from 7-4.0 Å;
  - XIII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of aspartic acid 48 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said aspartic acid 48 ranges from 2.7-4.0 Å;
- 30 XIV. a hydrophilic group that interacts with the backbone amide carbonyl group of asparagine 44 such that the distance between the centroid of said-hydrophilic group and the amide carbonyl group of said asparagine 44 ranges from 2.7-4.0 Å;

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- XV. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;
- XVI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;
  - XVII. a hydrophobic group that reaches a proximity interacts with the side chain methylene groups of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;
- XVIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of arginine 45 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said arginine 45 ranges from 2.7-4.0 Å;
- XIX. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of tyrosine 46 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said tyrosine 46 ranges from 2.7-4.0 Å;
- XX. a hydrophilic group that forms a hydrogen bond with the side chain amino group of lysine 41 such that the distance between the centroid of said hydrophilic group and the amino group of said lysine 41 ranges from 2.7-4.0 Å;
- XXI. a hydrophobic group that interacts with the side chainmethylene groups of lysine 41 such that the distance between the centroid

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of said hydrophilic group and the centroid of the methylene groups of said lysine 41 ranges from 4.4-5.1 Å;

- XXII. a hydrophobic group that interacts with the side chain methylene groups of leucine 88 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said leucine 8 ranges from 4.4-5.1 Å;
- XXIII. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of serine 118 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said serine 118 ranges from 2.7-4.0 Å;
  - XXIV. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of leucine 119 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said leucine 119 ranges from 2.7-4.0 Å;
- XXV. a hydrophilic group that forms a hydrogen bond with the one of the hydrogen atoms donated by the side chain amide nitrogen of glutamine 262 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glutamine 262 ranges from 2.7-4.0 Å;
- 25 XXVI. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide group nitrogen of glycine 259 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glycine 259 ranges from 2.7-4.0 Å;
  - XXVII. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the side chain guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

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XXVIII. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

XXIX. a hydrophobic group that interacts with the side chain methylene groups of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 254 ranges from 4.4-5.1 Å;

XXX. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

XXXI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

XXXII. a hydrophobic group that interacts with the side chain methylene groups of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;

XXXIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of aspartic acid 48 such that the distance between the centroid of said hydrophilic group and the backbone amide carbonyl group of said aspartic acid 48 ranges from 2.7-3.5 Å;

XXXIV. a hydrophobic group that interacts with the side chain atoms of methionine 258 such that the distance between the centroid of

said hydrophobic group and the centroid of the side chain of said methionine 258 ranges from 4.5-6.2 Å;

XXXV. a hydrophobic group that interacts with glycine 259 such that the distance between the centroid of said hydrophobic group and the centroid of the alpha-carbon atom of said glycine 259 ranges from 4.5-6.2 Å;

XXXVI. a hydrophobic group that interacts with phenylalanine 52 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic group of said phenylalanine 52 ranges from 4.1-9.1 Å; or

XXXVII.a hydrophobic group that interacts with methionine 258, glycine 259 and phenylalanine 52 being part of a hydrophobic pocket such that the distance between the centroid of said hydrophobic group and (i) the centroid of the side chain of said methionine 258 ranges from 4.1-7.2 Å, (ii) the centroid of said glycine 259 ranges from 4.7-7.7 Å, and (iii) the centroid of the side chain of said phenylalanine 52 ranges from 4.1-9.1 Å;

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In another embodiment, the invention provides a method of inhibiting at least one PTPase selected from the group consisting of PTP1B, TC-PTP andother PTPase that are structurally similar to PTP1B comprising exposing said PTPase to a compound that fits spatially into the active site of said PTPase and the vicinity therof, said compound comprising the following features and moieties:

I. (a) a phosphate isostere which forms a salt bridge to the guanidinium group of arginine 221 and interacts with a hydrogen atom donated by the backbone amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the phosphate isostere group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 backbone amide nitrogen ranges from 3.5-4.2 Å, and (III) said glycine 220 backbone amide nitrogen ranges from 2.7-3.5 Å; or (b) an oxalylamide which forms a salt bridge to the guanidinium

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group of arginine 221 and forms a hydrogen bond with a hydrogen atom donated by the amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the carboxylic acid group of said oxalylamide group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 amide nitrogen ranges from 3.5-4.2 Å and the distance between the amide carbonyl group of said oxalylamide group and the said glycine 220 amide nitrogen ranges from 2.7-3.5 Å; and

10 II. (a) a carboxylic acid group or (b) a carboxylic acid isostere group selected from the following 5-membered heterocycles

wherein said acid or acid isostere group forms a salt bridge to the side chain amino group of lysine 120 such that the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said lysine 120 ranges from 3.4-4.1 Å; and

III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å; and

one or more of the following features IV and V:

- IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 4.4-5.1 Å; and/or

- VI. an amino group which forms a salt bridge to the side chain carboxylic acid group of aspartic acid 48 such that the distance between the nitrogen atom of said amino group and the centroid of said side chain carboxylic acid group of aspartic acid 48 ranges from 3.4-4.1 Å; and
- VII. two oxygen atoms which form hydrogen bonds via a water molecule to the side chain carboxylic acid group of aspartic acid 48 such that the distance between each of the two oxygen atoms and the centroid of said water molecule ranges from 2.5-3.6 Å and that the distance between said water molecule and the centroid of said side chain carboxylic acid group of aspartic acid 48 ranges from 2.5-3.6 Å and that the distance between said two oxygen atoms ranges from 2.5-3.0 Å; and

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VIII. a hydrophobic group that interacts with the side chain methylene groups of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the methylene groups of said tyrosine 46 ranges from 4.4-5.1 Å;

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IX. a hydrophilic group that forms a salt bridge with aspartic acid 181 such that the distance between the centroid of said hydrophilic group and the centroid of the carboxylic acid of said aspartic acid 181 ranges from 4.4-5.1 Å;

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X. a hydrophobic group that interacts with tyrosine 46 and the methylene side chain atoms of arginine 47 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.7-5.2 Å and the centroid of the methylene side chain atoms of said arginine 47 ranges from 4.5-5.5 Å;

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XI. a-hydrophilic group that forms a hydrogen bond with the one or more hydrogen atoms donated by the guanidinium group of arginine 47 such that the distance between the centroid of said hydrophilic

group and the guanidinium group of said arginine 47 ranges from 2.7-3.5 Å;

- XII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of arginine 47 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said arginine 47 ranges from 2.7-4.0 Å;
- XIII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of aspartic acid 48 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said aspartic acid 48 ranges from 2.7-4.0 Å;
  - XIV. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of asparagine 44 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said asparagine 44 ranges from 2.7-4.0 Å;
- 20 XV. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

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XVI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

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XVII. a hydrophobic group that interacts with the side chain methylene groups of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;

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- XVIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of arginine 45 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said arginine 45 ranges from 2.7-4.0 Å;
- XIX. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of tyrosine 46 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said tyrosine 46 ranges from 2.7-4.0 Å;
- XX. a hydrophilic group that forms a hydrogen bond with the side chain amino group of lysine 41 such that the distance between the centroid of said hydrophilic group and the amino group of said lysine 41 ranges from 2.7-4.0 Å;
- XXI. a hydrophobic group that interacts with the side chain methylene groups of lysine 41 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said lysine 41 ranges from 4.4-5.1 Å;
- XXII. a hydrophobic group that interacts with the side chain methylene groups of leucine 88 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said leucine 8 ranges from 4.4-5.1 Å;
- XXIII. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of serine 118 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said serine 118 ranges from 2.7-4.0 Å;
- AXXIV.—a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of leucine 119 such that the distance

between the centroid of said hydrophilic group and the amide carbonyl group of said leucine 119 ranges from 2.7-4.0 Å;

XXV. a hydrophilic group that forms a hydrogen bond with the one of the hydrogen atoms donated by the side chain amide nitrogen of glutamine 262 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glutamine 262 ranges from 2.7-4.0 Å;

XXVI. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide group nitrogen of glycine 259 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glycine 259 ranges from 2.7-4.0 Å;

XXVII. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the side chain guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

XXVIII. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

XXIX. a hydrophobic group that interacts with the side chain methylene groups of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 254 ranges from 4.4-5.1 Å;

XXX. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and

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the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

XXXI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

XXXII. a hydrophobic group that interacts with the side chain methylene groups of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;

XXXIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of aspartic acid 48 such that the distance between the centroid of said hydrophilic group and the backbone amide carbonyl group of said aspartic acid 48 ranges from 2.7-3.5 Å;

XXXIV. a hydrophobic group that interacts with the side chain atoms of methionine 258 such that the distance between the centroid of said hydrophobic group and the centroid of the side chain of said methionine 258 ranges from 4.5-6.2 Å;

XXXV. a hydrophobic group that interacts with glycine 259 such that the distance between the centroid of said hydrophobic group and the centroid of the alpha-carbon atom of said glycine 259 ranges from 4.5-6.2 Å;

SXXVI. a hydrophobic group that interacts with phenylalanine 52 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic group of said phenylalanine 52 ranges

—from 4-1-9.1 Å; or

XXXVII.a hydrophobic group that interacts with methionine 258, glycine 259 and phenylalanine 52 being part of a hydrophobic pocket such that the distance between the centroid of said hydrophobic group and the centroid of the side chain of said methionine 258 ranges from 4.1-7.2 Å, the centroid of said glycine 259 ranges from 4.7-7.7 Å, and the centroid of the side chain of said phenylalanine 52 ranges from 4.1-9.1 Å;

In yet another embodiment, the invention provides a method of inhibiting a PTPase selected from the group consisting of PTP1B, TC-PTP and other PTPases that are structurally similar to PTP1B comprising exposing said PTPase to a compound that fits spatially into the active site of said PTPase and the vicinity thereof, said compound comprising the following features and moieties:

(a) a phosphate isostere which forms a salt bridge to the quanidinium group of arginine 221 and interacts with a hydrogen atom donated by the backbone amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the phosphate isostere group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 backbone amide nitrogen ranges from 3.5-4.2 Å, and (III) said glycine 220 backbone amide nitrogen ranges from 2.7-3.5 Å; or (b) an oxalylamide which forms a salt bridge to the guanidinium group of arginine 221 and forms a hydrogen bond with a hydrogen atom donated by the amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the carboxylic acid group of said oxalylamide group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 amide nitrogen ranges from 3.5-4.2 Å and the distance between the amide carbonyl group of said oxalylamide group and the said glycine 220 amide nitrogen ranges from 2.7-3.5 Å; and

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II. (a) a carboxylic acid group or (b) a carboxylic acid isostere group selected from the following 5-membered heterocycles

wherein said acid or said isostere group forms a salt bridge to the side chain amino group of lysine 120 such that the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said Lysine 120 ranges from 3.4-4.1 Å; and

III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å; and

at lest one of the following features IV and V:

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IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 3.55.1 Å; and/or

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V. a hydrophobic group that interacts with the imidazole ring of histidine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said histidine 182 is 4.4-6.5 Å; and one or more of the following features VI-XXXVII

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VI. two oxygen atoms which form hydrogen bonds via a water molecule to the side chain carboxylic acid group of aspartic acid 48 such that the distance between each of the two oxygen atoms and the centroid of said water molecule ranges from 2.5-3.6 Å and that the distance between said water molecule and the centroid of said side chain carboxylic acid group of aspartic acid 48 ranges from 2.5-3.6 Å and that the distance between said two oxygen atoms ranges from 2.5-3.0 Å;

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VII. an amino group which forms a salt bridge to the side chain carboxylic acid group of aspartic acid 48 such that the distance between the nitrogen atom of said amino group and the centroid of said side chain carboxylic acid group of aspartic acid 48 is 3.4-4.1 Å;

VIII. a hydrophobic group that interacts with the side chain methylene groups of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the methylene groups of said tyrosine 46 ranges from 4.4-5.1 Å;

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IX. a hydrophilic group that forms a hydrogen bond with aspartic acid 181 such that the distance between the centroid of said hydrophilic group and the centroid of the carboxylic acid of said aspartic acid 181 ranges from 4.4-5.1 Å;

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- X. a hydrophobic group that interacts with tyrosine 46 and the methylene side chain atoms of arginine 47 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.7-5.2 Å and the centroid of the methylene side chain atoms of said arginine 47 ranges from 4.5-5.5 Å;
- XI. a hydrophilic group that forms a hydrogen bond with the one or more hydrogen atoms donated by the guanidinium group of arginine 47 such that the distance between the centroid of said hydrophilic group and the guanidinium group of said arginine 47 ranges from 2.7-3.5 Å;
- XII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of arginine 47 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said arginine 47 ranges from 2.7-4.0 Å;
- XIII. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide nitrogen of aspartic acid 48 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said aspartic acid 48 ranges from 2.7-4.0

XIV. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of asparagine 44 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said asparagine 44 ranges from 2.7-4.0 Å;

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- XV. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;
- XVI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;
- XVII. a hydrophobic group that interacts with the side chain methylene groups of arginine 45 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;
- XVIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of arginine 45 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said arginine 45 ranges from 2.7-4.0 Å;
- XIX. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of tyrosine 46 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said tyrosine 46 ranges from 2.7-4.0 Å;
- a hydrophilic group that forms a hydrogen bond with the side chain amino group of lysine 41 such that the distance between the

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centroid of said hydrophilic group and the amino group of said lysine 41 ranges from 2.7-4.0 Å;

- XXI. a hydrophobic group that interacts with the side chain methylene groups of lysine 41 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said lysine 41 ranges from 4.4-5.1 Å;
- XXII. a hydrophobic group that interacts with the side chain methylene groups of leucine 88 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said leucine 8 ranges from 4.4-5.1 Å;
- XXIII. a hydrophilic group that forms a hydrogen bond with the side chain hydroxy group of serine 118 such that the distance between the centroid of said hydrophilic group and the hydroxy group of said serine 118 ranges from 2.7-4.0 Å;
  - XXIV. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of leucine 119 such that the distance between the centroid of said hydrophilic group and the amide carbonyl group of said leucine 119 ranges from 2.7-4.0 Å;
- XXV. a hydrophilic group that forms a hydrogen bond with the one of the hydrogen atoms donated by the side chain amide nitrogen of glutamine 262 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glutamine 262 ranges from 2.7-4.0 Å;
  - XXVI. a hydrophilic group that forms a hydrogen bond with the hydrogen atom donated by the backbone amide group nitrogen of glycine 259 such that the distance between the centroid of said hydrophilic group and the amide nitrogen group of said glycine 259 ranges from 2.7-4.0 Å;

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XXVII. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the side chain guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

XXVIII. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 254 ranges from 2.7-4.0 Å;

XXIX. a hydrophobic group that interacts with the side chain methylene groups of arginine 254 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 254 ranges from 4.4-5.1 Å;

XXX. a hydrophilic group that forms a hydrogen bond with one or more hydrogen atoms donated by the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å:

XXXI. a hydrophilic group that forms a salt bridge with the guanidinium group of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the guanidinium group of said arginine 24 ranges from 2.7-4.0 Å;

XXXII. a hydrophobic group that interacts with the side chain methylene groups of arginine 24 such that the distance between the centroid of said hydrophilic group and the centroid of the methylene groups of said arginine 24 ranges from 4.4-5.1 Å;

XXXIII. a hydrophilic group that forms a hydrogen bond with the backbone amide carbonyl group of aspartic acid 48 such that the distance

between the centroid of said hydrophilic group and the backbone amide carbonyl group of said aspartic acid 48 ranges from 2.7-3.5 Å;

XXXIV. a hydrophobic group that interacts with the side chain atoms of methionine 258 such that the distance between the centroid of said hydrophobic group and the centroid of the side chain of said methionine 258 ranges from 4.5-6.2 Å;

XXXV. a hydrophobic group that interacts with glycine 259 such that the distance between the centroid of said hydrophobic group and the centroid of the alpha-carbon atom of said glycine 259 ranges from 4.5-6.2 Å:

XXXVI. a hydrophobic group that interacts with phenylalanine 52 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic group of said phenylalanine 52 ranges from 4.1-9.1 Å; or

XXXVII.a hydrophobic group that interacts with methionine 258,
20 glycine 259 and phenylalanine 52 being part of a hydrophobic pocket such
that the distance between the centroid of said hydrophobic group and the
centroid of the side chain of said methionine 258 ranges from 4.1-7.2 Å,
the centroid of said glycine 259 is 4.7-7.7 Å, and the centroid of the side
chain of said phenylalanine 52 ranges from 4.1-9.1 Å;

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Further provided is a method of inhibiting at least one PTPase selected from the group consisting of Protein Tyrosine Phosphatase 1B (PTP1B) and/or T-Cell Protein Tyrosine Phosphatase which (TC-PTP) and/or other PTPases that are structurally similar to PTP1B comprising exposing said PTPase to a compound that fits spatially into the active site of said PTPase and the vicinity thereof, said compound comprising:

I. (a) a phosphate isostere which forms a salt bridge to the guanidinium group of arginine 221 and forms a hydrogen bond with a

hydrogen atom donated by the backbone amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the phosphate isostere group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 backbone amide nitrogen ranges from 3.5-4.2 Å, and (III) said glycine 220 backbone amide nitrogen ranges from 2.7-3.5 Å; or (b) an oxalylamide which forms a salt bridge to the guanidinium group of arginine 221 and forms a hydrogen bond with a hydrogen atom donated by the amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the carboxylic acid group of said oxalylamide group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 amide nitrogen ranges from 3.5-4.2 Å and the distance between the amide carbonyl group of said oxalylamide group and the said glycine 220 amide nitrogen ranges from 2.7-3.5 Å; and

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II. (a) a carboxylic acid group or (b) acarboxylic acid isostere group selected from the following 5-membered heterocycles

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wherein said acid or isostere group forms a salt bridge to the side chain amino group of lysine 120 such that the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said lysine 120 ranges from 3.4-4.1 Å; and

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III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å; and at least one of the following features IV and V:

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IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 4.4-5.1 Å; or

V. a hydrophobic group that interacts with the imidazole ring of histidine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said histidine 182 ranges from 4.4-6.5 Å.

In another specific embodiment, the invention provides a method of inhibiting at least one PTPase selected from the group consisting of Protein Tyrosine Phosphatase 1B (PTP1B), T-Cell Protein Tyrosine Phosphatase (TC-PTP) and other PTPases that are structurally similar to PTP1B which comprises exposing said PTPase to a compound that fits spatially into the active site of said PTPase and the vicinity thereof, said compound comprising:

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- I. a phosphate isostere which forms a salt bridge to the guanidinium group of arginine 221 and interacts with a hydrogen atom donated by the backbone amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the phosphate isostere group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 backbone amide nitrogen ranges from 3.5-4.2 Å, and (III) said glycine 220 backbone amide nitrogen ranges from 2.7-3.5 Å; and
- 25 II. (a) a carboxylic acid group or (b) a carboxylic acid isostere group selected from the following 5-membered heterocycles

wherein said acid or isostere group forms a salt bridge to the side chain amino group of lysine 120 such that the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said lysine 120 ranges from 3.4-4.1-Å; and

III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å; and

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IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 4.4-5.1 Å; or

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V. a hydrophobic group that interacts with the imidazole ring of histidine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said histidine 182 ranges from 4.4-6.5 Å;

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wherein the distance between the centroid of the phosphate isostere and the centroid of (I) said carboxylic acid or carboxylic acid isostere ranges from 4.9-5.9 Å, (II) said amino group ranges from 8.0-14.0 Å and between the centroid of said carboxylic acid or carboxylic acid isostere and said amino group ranges from 4.8-5.8 Å or

wherein the distance between the centroid of the phosphate isostere and the centroid of (I) said carboxylic acid or carboxylic acid isostere ranges from 4.9-5.9 Å, (II) said oxygen atoms are ranges from 8.0-14.0 Å and between the centroid of said carboxylic acid or carboxylic acid isostere and said oxygen atoms are ranges from 5.0-7.9 Å.

The invention further provides a method of inhibiting at least one PTPase selected from the group consisting of Protein Tyrosine Phosphatase 1B (PTP1B), T-Cell Protein Tyrosine Phosphatase (TC-PTP) and other PTPases that are structurally similar to PTP1B which comprises exposing said PTPase to a compound that fits spatially into the active site of said PTPase and the vicinity thereof, said compound comprising:

- I. an oxalylamide which forms a salt bridge to the guanidinium group of arginine 221 and forms a hydrogen bond with a hydrogen atom donated by the amide nitrogens of arginine 221 and glycine 220 such that the distance between the centroid of the carboxylic acid group of said oxalylamide group and (I) the centroid of said guanidinium group ranges from 3.50-4.20 Å, (II) said arginine 221 amide nitrogen ranges from 3.5-4.2 Å and the distance between the amide carbonyl group of said oxalylamide group and the said glycine 220 amide nitrogen ranges from 2.7-3.5 Å; and
  - II. (a) a carboxylic acid group or (b) a carboxylic acid isostere group selected from the following 5-membered heterocycles

wherein said acid or isostere group forms a salt bridge to the side chain amino group of lysine 120 such that the distance between the centroid of said carboxylic acid or carboxylic acid isostere and the side chain nitrogen atom of said lysine 120 ranges from 3.4-4.1 Å; and

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III. a hydrophobic group that interacts with the aromatic ring of tyrosine 46 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said tyrosine 46 ranges from 4.4-5.1 Å; and

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IV. a hydrophobic group that interacts with the aromatic ring of phenylalanine 182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said phenylalanine 182 ranges from 4.4-5.1 Å; or

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V. a hydrophobic group that interacts with the imidazole ring of-histidine-182 such that the distance between the centroid of said hydrophobic group and the centroid of the aromatic ring of said histidine 182 ranges from 4.4-6.5 Å; and

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wherein the distance between the centroid of the carboxylic acid group of said oxalylamide group and the centroid of (I) said carboxylic acid or carboxylic acid isostere ranges from 4.9-5.9 Å, (II) said amino group ranges from 8.0-14.0 Å and between the centroid of said carboxylic acid or carboxylic acid isostere and said amino group ranges from 4.8-5.8 Å or

wherein the distance between the centroid of the carboxylic acid group of said oxalylamide group and the centroid of (I) said carboxylic acid or carboxylic acid isostere ranges from 4.9-5.9 Å, (II) said oxygen atoms are ranges from 8.0-14.0 Å and between the centroid of said carboxylic acid or carboxylic acid isostere and said oxygen atoms are ranges from 5.0-7.9 Å.

The hydrophobic groups that interact with tyrosine 46 and phenylalanine/histidine 182 include, but are not limited to, alkyl and aryl groups. These hydrophobic groups include cyclohexyl, phenyl, naphthyl, thiophenyl, pyrrolyl and furanyl. The hydrophobic groups that interact with one or more of the tyrosine 46 and the arginines 24, 45, 47, and 254 include, but are not limited to, alkyl and aryl groups. These hydrophobic groups include cyclohexyl, phenyl, naphthyl,thiophenyl, pyrrolyl and furanyl, optionally substitutedThe hydrophobic groups that interact with methionine 258, glycine 259 and phenylalanine 52 include, but are not limited to, alkyl and aryl groups groups. These aryl groups include phenyl, thiophenyl, pyrrolyl, furanyl, C<sub>1</sub>-C<sub>6</sub>alkyl and arylC<sub>1</sub>-C<sub>6</sub>alkyl which are defined hereinbelow.

The hydrophilic groups that interact with the hydrogen atom donated by the side chain amide nitrogen of arginine 47, aspartic acid 48, leucine 119, glycine 259, lysine 41, lysine 120, the side chain amide hydrogen atom donated by glutamine 262, the hydrogen atoms donated by the guanidinium group of arginine 254, arginine 45 or arginine 24 include, but are not limited to, hydroxy,  $C_1$ - $C_6$ alkyloxy, aminocarbonyl, oxo, SO,  $SO_2$ ,

SONH<sub>2</sub>, SO<sub>2</sub>NH<sub>2</sub>, SO<sub>2</sub>NHCF<sub>3</sub>, COOH or a group selected from the following 5-membered heterocycles

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The hydrophilic groups that interact with the side chain amide carbonyl group of asparagine 44, arginine 45 or aspartic acid 48 include, but are not limited to, amino, aminocarbonyl, hydroxy, SONH<sub>2</sub>, SO<sub>2</sub>NH<sub>2</sub>, or SO<sub>2</sub>NHCF<sub>3</sub>.

The hydrophilic groups that interact with the side chain carboxylic acid group of aspartic acid 181 include, but are not limited to, amino, aminocarbonyl, hydroxy, C<sub>1</sub>-C<sub>6</sub>alkyloxy, SONH<sub>2</sub>, SO<sub>2</sub>NH<sub>2</sub>. The hydrophilic groups that interact with the side chain hydroxy group of serine 118 include, but are not limited to, aminocarbonyl, hydroxy, C<sub>1</sub>-C<sub>6</sub>alkyloxy, SONH<sub>2</sub>, SO<sub>2</sub>NH<sub>2</sub>.

# Unique structural elements in PTP1B

To identify unique residues or combinations of residues of PTP1B that could be utilised as points of interaction by selective inhibitors, alignment of the primary sequences of the catalytic domains of approximately 105 known vertebrate PTPases (Andersen, J.N. *et al.*, (1999) in preparation) was done (Table 1, below). Using the crystal structure of PTP1B (Andersen, H.S. *et al.* (1999) *J. Biol. Chem.* **275**:-7107-7108 (2000);Barford, D., *et al. Science* **263**:1397-1404 (1994)), unique combinations of residues in the active site pocket or in its vicinity were

identified, i.e. in a distance (3-5.5 Å) that would allow simultaneous binding to the active site and these residues, while still retaining a low molecular weight (for example, below 700 dalton). In particular, the combination of 4 residues seems unique for the PTP1B family: arginine 47, aspartic acid 48, methionine 258, and glycine 259, arginine 47 and aspartic acid 48 contribute significantly to the binding of peptide substrates in PTP1B (Jia, Z.C., et al., Science 268:1754-1758 (1995)). A comparison of these regions in representative members of 14 PTP families, indicates that in particular residue 48 is an attractive binding element for selective PTP1B ligands since this residues is an aspartic acid in PTP1B and an asparagine in many other PTPases. Aspartic acid 48 is well-defined in the published PTP1B structures ((Puius, Y.A. et al. Proc. Natl. Acad. Sci. USA 94:13420-13420 (1997)), (Pannifer, A.D.B., et al., J. Biol. Chem. 273:10454-10462 (1998)) and it is believed to play an 15 important role in positioning substrates correctly relative to the active site (Sarmiento, M., et al., J. Biol. Chem. 273: 26368-26374 (1998)).

#### 20 **Table 1**

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Non-limiting examples of selected amino acid residues at positions in the vicinity of the active site (single letter code – PTP1B numbering)

Qe due	A A	b K	R	i de	St afest	E E	3 2	A A	R		i de	ge of	Š	a de
47	R	K	K	. R	K	. K	Р	. N	· v	P.	Α	G	V	1
48	. D	N	N	E	D.	. T	l D	N	N,	Ň	. N	. N	D	N
258	М	S	, H	М	A	G	Р	V	С	Р	. N	V	С	. N
259	G	G	G	F	M	G	G	Н	· Q	0	Y	N	L	Y

### **Optimization for potency**

The key structural features of 2-(oxalyl-amino)-benzoic acid (OBA) are
the twocarboxy groups respectively bound - directly and through a carbonylamino group - to an aromatic ring. Replacement of the phenyl ring

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in OBA by thiophene, resulted in compounds with little difference in potency between the regioisomer 2-aminothiophene and 3-aminothiophene.

Previous studies have shown that phenyl-based phosphonate inhibitors have little affinity for PTP1B, while addition of a second phenyl ring (e.g. [(1,1-difluoro-1-naphthalenyl)-methyl]phosphonic significantly acid) increased the potency (Burke, T.R. et al., Biochemistry 35:15989-15996 (1996)). The enhanced potency of the naphthalene ring system is due to extensive hydrophobic interactions with the side chains of tyrosine 46, valine 49, phenylalanine 182, alanine 217 and isoleucine 219. Similarly, 3-(oxalyl-amino)-naphthalene-2-carboxylic acid interacts with the same residues. It was reasoned that a saturated ring fused to 2-(oxalyl-amino)thiophene-3-carboxylic acid (2-OTA) and/or 3-(oxalyl-amino)-thiophene-2carboxylic acid (3-OTA) would serve a similar function and increase the potency. Further, the proposed binding mode of such a compound should bring the saturated ring in close proximity to residues arginine 47 and aspartic acid 48. Introducing a basic nitrogen or polar changes in this saturated ring would allow further interactions with the side chains or backbone amides of arginine 47 and aspartic acid 48. In accordance with the above alignment studies, we anticipated that selectivity for PTP1B and other PTPases with an aspartic acid in position 48 could be obtained by specifically addressing this area of the enzyme.

Consequently, 2-(oxalyl-amino)-4,5,6,7-tetrahydro-benzo[b]thiophene-3-carboxylic acid (2-OBTA) was synthesised and its potency analysed against a set of PTPases. Table II shows that 2-OBTA is about 10-fold more potent against PTP1B than compounds 3-OTA and 2-OTA and 3-fold more potent than OBA when tested at pH 5.5 (the pH optimum for PTP1B using pNPP as substrate). Further, the inhibitory profile against this set of PTPases is almost the same as that of 3-(oxalyl-amino)-naphthalene-2-carboxylic acid. Thus, although 2-OBTA retains the features of a general PTP inhibitor, it already shows some selectivity for PTP1B. These results clearly indicate that 2-OBTA spatially fits in this region of PTP1B. Various substitutions in the saturated ring of 2-OBTA

were found to influence the binding affinities for different PTPases (not shown).

Table 2 K<sub>i</sub> values (μM) – pH 5.5

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	OBA	3-OTA	2-OTA	2-OBTA	2-OTPyA	2-OTPA
PTP1B	20	61	62	5.7	0.3	15
SHP-1	530	>2000	60	120	900	350
PTPα D1	700	500	1700	300	>2000	270
ΡΤΡε D1	125	350	590	45	600	20
ΡΤΡβ	32	160	18	14	150	12
CD45 D1D2	160	250	70	40	110	50
LAR D1D2	>2000	>2000	>2000	400	>2000	360

As indicated above, in comparison with OBA, 2-OBTA showed an approximately 3-fold increase in affinity for most PTPases. It was hypothesised that the saturated ring of 2-OBTA would occupy almost the same position as the distal ring of 3-(oxalyl-amino)-naphthalene-2-carboxylic acid, which was previously shown to bind in the proximity of arginine 47 and aspartic acid 48. Therefore, as expected, there was no apparent change in selectivity in accordance with the notion that the saturated ring makes hydrophobic contact with conserved residues such as tyrosine 46, alanine 217, valine/isoleucine219 and isoleucine/valine 49 (PTP1B numbering).

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### **Optimization for selectivity**

The combination of arginine 47 and aspartic acid 48 offers a rather unique, selective ligand-binding region in PTP1B. The side chains of both residues are charged at neutral pH and are therefore sutiable for salt bridge formation. Introducing a positive charge in 2-OBTA that could form a salt bridge with aspartic acid 48, would not only increase the potency of 20BTA against PTP1B but also — due to repulsive forces between the positive ligand charge and the asparagine side chain found in many other PTPases — decrease the affinity of 20BTA for these PTPases.

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Three side chain rotamer conformations are normally defined for an aspartic acid residue (rota 1: 47.7%, rota 2: 33.6% and rota 3: 15.9%). In the published X-ray structures of PTP1B, two rotamers have been described, rota 1 and 3. The rota 3 conformation is stabilised by an internal hydrogen bond between the side chain and main chain amide with the side chain bending towards the active site pocket. Further, rota 3 seems to be the preferred rotamer for aspartic acid 48. The rota 1 conformation has only been found in four of the eleven published X-ray structures, and in three of these cases the rota 1 position is necessitated due to ligand occupancy. The aspartic acid 48 rota 1 conformation is pointing away from the active site pocket. Thus, rota 3 was found both in the apo-enzyme and in PTP1B complexed with peptide ligands that seem to stabilize this conformation. Further, we have recently co-crystallized PTP1B with OBA and 3 derivatives and found aspartic acid 48 in the rota 3 position in all structures (Andersen, H.S. et al. J. Biol. Chem. 275, 7101-7108 (2000)). Based on these observations, it was hypothesized that introduction of a basic nitrogen in the saturated ring in 2-OBTA would be sufficiently close to aspartic acid 48 to allow the formation of a salt bridge. A recent survey of 322 unrelated proteins has shown that aspartic acid and asparagine residues have a strong tendency to form hydrogen bonds with neighboring backbone amides and in both cases with a significant preference for internal hydrogen bonds.

Assuming that asparagine 48 of other PTPases, e.g.  $PTP\alpha$ , forms an internal hydrogen bond similar to that observed for aspartic acid 48 in PTP1B, the side chain amide of the asparagine with its positive dipole would be in an unfavourable position to the proposed basic nitrogen and thus cause repulsion.

2-(Oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid (2-OTPyA) - containing a positively charged tetrahydropyridine ring (p $K_a$  > 10) - was synthesised in order to test the foregoing hypothesis. In agreement with the predictions, the affinity for PTP1B was increased about 20-fold without any significant increase in molecular weight (Table 2). Further, this compound showed an almost astonishing selectivity for

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PTP1B *versus* all other PTPases tested. Again, this is in agreement with the hypothesis that repulsive forces between the basic nitrogen in 2-OTPyA and the positive dipole of the asparagine side chain decrease the potency against other PTPases. CD45, which also contains an aspartic acid in position 48, is a noticeable exception showing only a 2-fold decrease. It is speculated that the preferred rotamer of aspartic acid 48 in CD45 is the rota 1 conformation, which is too far away for salt bridge formation with 2-OTPyA. In addition, CD45 contains a valine in position 47, which may not have the same influence on aspartic acid 48 as an arginine.

2-(Oxalyl-amino)-4,7-dihydro-thieno[2,3-c]pyran-3-carboxylic acid (2-OTPA) - containing a negative dipole in the dihydropyran ring - was synthesised. In agreement with the predictions, the affinity for PTP1B was decreased about 2.5-fold compared to 2-OBTA without any significant increase in molecular weight (Table 2).

Table A (at the end of the specification) discloses the protein coordinates of PTP1B complexed with 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid (2-OTPA) and in Figure 1 is the active site of PTP1B complexed with 2-OTPA shown.

Optimization for potency towards Arginine 47 and Aspartic acid 48
Using further the combination of the 4 unique residues for the PTP1B
family: arginine 47, aspartic acid 48, methionine 258, and glycine 259 it was hypothesised that an increase in potency could be obtained by introduction of a hydrogen-bond acceptor side chain that could form one or more hydrogen bonds with the main chain amides of arginine 47 and aspartic acid 48, would increase the potency against PTP1B.

5-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid (5-HTPyA) (Example 52) - still containing a positively charged tetrahydropyridine ring and three hydrogen-bond acceptors (oxygen atoms) - was synthesised. In

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agreement with the predictions, the affinity for PTP1B was increased about 13- fold compared to 2-OTPyA.

Changing the positively charged nitrogen atom with a non charged oxygen atom and still addressing the main chain amides of arginine 47 and

5 aspartic acid 48, it was hypothesised that an increase in general potency could be obtained. Thus, 5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (5-HTPA) (Example 4) - containing a non charged dihydropyran ring and three hydrogen-bond acceptor oxygen atoms - was synthesised. In

10 agreement with the predictions, only the general potency was increased compared to 2-OTPyA as shown in Table 3.

# 15 Selectivity via steric hindrance

Referring again to the combination of the 4 residues unique for the PTP1B family: arginine 47, aspartic acid 48, methionine 258, and glycine 259, but this time more specifically to the combination of methionine 258 and glycine 259, which form part of a hydrophobic pocket in PTP1B in contrast to most other PTPases where the pocket is filled out: PTPa: cysteine 258glutamine 259; PTP β: valine 258-histidine 259; PTP-LAR: asparagine 258-tyrosine 259; and CD45: cysteine 258-leucine 259 (PTP1B) numbering), it was hypothesised that an increase in potency and selectivity could be obtained by introduction of a hydrophobic side chain that could form hydrophobic interactions to glycine 259 and to the side chain of methionine 258 and at the same time take part in repulsion-/steric hindrance with the same residues in other PTPases. Thus, 7-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (7-MOTPA) (Example 26) -containing a hydrophobic 1,3-dihydro-isoindol side chain -- was synthesised. In agreement with the predictions, both affinity and selectivity for PTP1B was increased as shown in Table 3 compared to 2-OTPA.

Table 3

K<sub>i</sub> values (μM) – pH 7

	2-OTPA	7-MOTPA	5-HTPA
PTP1B	63	1.2	1.9
PTPα D1	1100	620	93
PTPε D1	290	330	11
ΡΤΡβ	17	8:9	1.1
CD45 D1D2	960	380	130

Table B (at the end of the specification) discloses the protein coordinates of PTP1B complexed with 7-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (7-MOTPA) (Example 26), and Figure 2 shows the crystal structure of the active site of PTP1B complexed with 7-MOTPA.

Table C (at the end of the specification) discloses the protein coordinates of PTP1B complexed with 5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (5-HTPA) (Example 4), and Figure 3 shows the crystal structure of the active site of PTP1B complexed with 5-HTPA.

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Table D (at the end of the specification) discloses the protein coordinates of PTP1B complexed with 2-(oxaly1-amino)-7-(1,1, 3-trioxo-1*H* -benzo [d] isothiazol-3-yloxomethyl)-4,7-dihydro-5*H*, thieno [2,3-c] pyran-3- carboxylic acid (example 54), including key water molecules. Figure 2 is the active site with selected water molecules shown.

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Specific interactions of certain inhibitors of the present invention at the active site of PTP1B are detailed below.

The carboxy group of the oxamicN acid of 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid is positioned 2.9-3.0 Å from the guanidinium group of arginine 221 forming a salt bridge, as well as a hydrogen-bond with the main chain-amide-of-arginine 221 and serine 216, and the carbonyl forms a hydrogen bond with the main chain amide

30 of glycine 220. The carboxy group in the 3 position is positioned 2.8 Å

from lysine 120 forming a salt bridge. The tetrahydro-thieno[2,3-c]pyridine ring forms hydrophobic interactions with phenylalanine 182, tyrosine 46, valine 49, alanine 217 and isoleucine 219. The basic nitrogen in the tetrahydro-thieno[2,3-c]pyridine ring is positioned 2.8 Å from the carboxy group of aspartic acid 48 forming a salt bridge.

The carboxy group of the oxamic acid of 7-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-thieno[2,3-c]pyran-3-carboxylic acid (Example 26) is positioned 2.9-3.0 Å from the guanidinium group of arginine 221 forming a salt bridge, as well as a hydrogen bond with the main chain amide of arginine 221 and serine 216, and the carbonyl forms a hydrogen bond with the main chain amide of glycine 220. The carboxy group in the 3 position is positioned 2.8 Å from lysine 120 forming a salt bridge. The dihydro-thieno[2,3-c]pyran ring forms hydrophobic interactions with phenylalanine 182, tyrosine 46, valine 49, alanine 217 and isoleucine 219. The phenyl ring of the isoindol ring forms a hydrophobic interaction with the side chain methylene atom of aspartic acid 48 and the 5-methoxy substituent forms hydrophobic interactions with the side chain atoms of methionine 258.

The carboxy group of the oxamic acid of 5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-thieno[2,3-c]pyran-3-carboxylic acid (Example 4) is positioned 2.9-3.0 Å from the guanidinium group of arginine 221 forming a salt bridge, as well as a hydrogen bond with the main chain amide of arginine 221 and serine 216, and the carbonyl forms a hydrogen bond with the main chain amide of glycine 220. The the carboxy group in the 3 position is positioned 2.7 Å from lysine 120 forming a salt bridge. The dihydro-thieno[2,3-c]pyran ring forms hydrophobic interactions with phenylalanine 182, tyrosine 46, valine 49, alanine 217 and isoleucine 219. The side chain methylene group at the 5 position of the thieno[2,3-c]pyran forms a hydrophobic interaction the side chain methylene group of aspartic acid 48. The phenyl ring of the isoindol ring forms a hydrophobic interaction with tyrosine 46 and both one of the oxo atoms and the hydroxy group at the isoindole forms hydrogen

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bonds respectively with the main chain amide of aspartic acid 48 and arginine 47.

To further substantiate the generality in using steric hindrance/steric fit to obtain selectivity for PTP1B, TC-PTP and structurally similar PTPases we also synthesized 7-(1,1-dioxo-1H-benzo[d]isothiazol-3-yloxymethyl)-2-(oxalyl-amino)-4,7-dihyd ro-5H-thieno[2,3-c]pyran-3-carboxylic acid ("Compound N"). The substitution was introduced in the 7-position to address the region defined 10 by residues 258 and 259. As indicated above, this part of PTP1B forms a hydrophobic pocket with direct access to the active site, whereas the same region is sterically hindered by more bulky side chains, in particular those corresponding to residue 259 in PTP1B. Compound N was synthesized with a substituent in the 7-position of 2-OTPA to sterically fit with this part of PTP1B and TC-PTP, but cause steric hindrance in other 15 PTPs.

To test directly, whether the above compound was addressing the proposed region of PTP1B, Compound N was subjected to detailed enzyme kinetic analyses using a set of wildtype (wt) and mutant PTPs. 20 Two enzymes, PTP $\alpha$  and PTPH1, were chosen as representatives for PTPs with bulky side chains in the 259 position. Using a combination of wt and PTP mutants it has previously been shown that Gln259 in PTP $\alpha$ , in addition to its direct effect, also indirectly influences the binding of inhibitors and substrates, most likely due to a negative influence on the 25 rotational freedom of the side chain of Gln262 (Peters et al., J. Biol. Chem. 275: 18201-18209 (2000)). As described above, selectivity can be obtained by introducing a basic nitrogen into 2-(oxalylamino)-4,5,6,7tetrahydrobenzo[b]thiophene-3-carboxylic acid that causes attraction in PTP1B due to salt bridge formation to Asp48 and repulsion against PTPs 30 with an asparagine in the 48 position, such as PTP $\alpha$ . To analyze if the current approach based on steric hindrance is generally applicable, it was decided to include a PTP with an aspartic acid in position 48. PTPH1, which like PTP1B is an intracellular enzyme with one domain only, was

selected for these studies. The results of these studies are shown below (Table 4).

Table 4

Enzyme	Ki values (μM) at pH 7.0
PTP1B wt	0.4
PTP1B G259Q	65
PTP1B G259M	55
PTPα wt	>500
PTPα Q259G	70
PTPH1 wt	55
PTPH1 M259G	12

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It appears that introduction of bulky side chains in the 259 position in PTP1B causes a very significant decrease in affinity for NNC 52-1153. Conversely, replacement of the bulky residues in PTP $\alpha$  and PTPH1 with a glycine increases the affinity. This clearly indicates that NNC 52-1153 addresses the 258-259 region of PTP1B.

Specificity against a broad set of PTPs –It was next analyzed if the side chain of NNC 52-1153 would cause the increased selectivity against other PTPs. NNC 52-1153 was tested against a set of 10 different wt PTP domains (Table 5). It appears from this table that a substantial increase in affinity for PTP1B and TC-PTP has been obtained, while at the same time introducing a very high degree of selectivity against many other PTPs

representing a broad spectrum of this class of enzymes (having Asp 48).

### 20 Table 5

Enzyme	Ki values (μM) at pH 7.0
PTP1B	0.4
TC-PTP	0.6
PTPH1	55
ΡΤΡα	700
ΡΤΡε	460
CD45	500

LAR	120
GLEPP1	150
ΡΤΡβ	15

To unequivocally determine the binding mode, x-ray co-crystallization studies of PTP1B and NNC 52-1153 were initiated. A well-suited electron density was identified in the active site pocket. The oxalylamino and ocarboxy groups show the exact same interaction with the PTP signature motif and salt bridge formation to Lys120 as described previously for 2-(oxalylamino)-benzoic acid and the thiophene-based derivatives. Significantly, the side chain of the ligand is positioned in close vicinity to residues 258 and 259. Several interaction points appear to be responsible for the observed significant increase in affinity for PTP1B. Thus, a long hydrogen bond seems to interact with one carbonyl of the ligand side chain. In addition, important van der Waals contacts are made between the aromatic ring of the ligand side chain and the side chain of Met248 and Cβ atom of Asp48.

As described above, we have utilized salt bridge formation to Asp48 to obtain potent and selective PTP1B inhibitors. In these structures, Asp48 was in the so-called rotamer ("rota") 3 position – pointing towards the active site. In contrast, the side chain of Asp48 is pushed away from the active site by the oxygen molecules in NNC 52-1153 (i.e. the rotamer 1 position). This allows a novel water molecule to form a bridge between the two oxygen molecules in the ligand and Asp48. This surprising observation can be used to design additional inhibitors of PTP1B.

The present invention encompasses, but is not limited to, compounds of the Formula 1 wherein n, m, X, Y,  $R_1$ ,  $R_2$ ,  $R_3$ ,  $R_4$ ,  $R_5$  and  $R_6$  are defined below;

## Formula 1

- In the above Formula 1
   n is 0, 1 or 2 (if m = 0 then n is 1 or 2);
   m is 0, 1 or 2 (if n = 0 then m is 1 or 2);
   X is S, O, NR<sub>8</sub>;
   Y is NR<sub>7</sub>, O, S, SO, SO<sub>2</sub>;
- 10 R<sub>1</sub> is hydrogen, COOR<sub>3</sub>, or selected from the following 5-membered heterocycles:

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R<sub>2</sub> is hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, hydroxy, NR<sub>9</sub>R<sub>10</sub>;

R<sub>3</sub> is hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyloxyC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyloxyarylC<sub>1</sub>-C<sub>6</sub>alkyl;

20 R<sub>4</sub>, R<sub>5</sub> and R<sub>6</sub> are independently hydrogen, trihalomethyl, C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, hydroxy, oxo, carboxy, carboxyC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkyloxy-

substituted:

carbonyl, aryloxycarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkyloxycarbonyl, C<sub>1</sub>-C<sub>6</sub>alkyloxy, C<sub>1</sub>- $C_6$ alkyloxy $C_1$ - $C_6$ alkyl, aryloxy, aryloxy  $C_1$ - $C_6$  alkyl, aryl $C_1$ - $C_6$ alkyloxy,  $arylC_1-C_6alkyloxyC_1-C_6alkyl$ , thio,  $C_1-C_6alkyl$ thio,  $C_1-C_6alkyl$ thio $C_1-C_6alkyl$ , arylthio, arylC<sub>1</sub>-C<sub>6</sub>alkylthio, arylC<sub>1</sub>-C<sub>6</sub>alkylthioC<sub>1</sub>-C<sub>6</sub>alkyl, NR<sub>9</sub>R<sub>10</sub>, C<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, di(arylC<sub>1</sub>-C<sub>6</sub>alkyl)aminoC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarboxy, C<sub>1</sub>-C<sub>6</sub>alkylcarboxyC<sub>1</sub>-C<sub>6</sub>-alkyl, arylcarboxy, arylcarboxyC<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy, arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxyC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-10 C<sub>6</sub>alkylcarbonylamino, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl-aminoC<sub>1</sub>-C<sub>6</sub>alkyl, -carbonyINR<sub>7</sub>C<sub>1</sub>-C<sub>6</sub>alkyICOR<sub>13</sub>, aryIC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl-amino, aryIC<sub>1</sub>-C<sub>6</sub>alkylcarbonylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, arylamino carbonylaminoC<sub>1</sub>-C<sub>6</sub> alkyl, arylaminoC<sub>1</sub>-C<sub>6</sub> alkyl, arylcarbonylamino C<sub>1</sub>-C<sub>6</sub> alkyl, CONR<sub>9</sub>R<sub>10</sub>, R<sub>8</sub>R<sub>9</sub>NC<sub>1</sub>-C<sub>6</sub> alkyl, or C<sub>1</sub>-C<sub>6</sub> alkyl-CONR<sub>9</sub>R<sub>10</sub> wherein the alkyl and aryl groups are optionally substituted and  $R_{13}$  is  $NR_9R_{10}$ , or  $C_1$ - $C_6$ alky $INR_9R_{10}$ ; 15 R<sub>7</sub> is hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, C<sub>1</sub>-C<sub>6</sub>alkyloxocarbonyl, arylcarbonyl, aryloxocarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl,  $arylC_1-C_6alkyloxocarbonyl, C_1-C_6alkyloarboxy, arylC_1-C_6alkyloarboxy,$ R<sub>9</sub>R<sub>10</sub>NcarbonylC<sub>1</sub>-C<sub>6</sub>alkyl wherein R<sub>9</sub> and R<sub>0</sub> are independently selected

25  $R_8$  is hydrogen,  $C_1$ - $C_6$ alkyl, aryl, aryl $C_1$ - $C_6$ alkyl,  $C_1$ - $C_6$ alkylcarbonyl, aryl $C_1$ - $C_6$ alkylcarbonyl,  $C_1$ - $C_6$ alkylcarboxy or aryl $C_1$ - $C_6$ alkylcarboxy wherein the alkyl and aryl groups are optionally substituted;

from hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl,

arylcarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, C<sub>1</sub>-C<sub>6</sub>alkylcarboxy or arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy, wherein the alkyl and aryl groups are optionally

R<sub>9</sub> and R<sub>10</sub> are independently selected from hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, arylCarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy or arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy wherein the alkyl and aryl groups are optionally substituted; or R<sub>9</sub> and R<sub>10</sub> are together with the nitrogen to which they are attached forming a saturated, partially saturated or aromatic cyclic, bicyclic or

tricyclic ring system containing from 3 to 14 carbon atoms and from 0 to 3 additional heteroatoms selected from nitrogen, oxygen or sulphur, the ring system can optionally be substituted with at least one  $C_1$ - $C_6$ alkyl, aryl, aryl $C_1$ - $C_6$ alkyl, hydroxy, oxo,  $C_1$ - $C_6$ alkyloxy, aryl $C_1$ - $C_6$ alkyloxy,  $C_1$ -

- C<sub>6</sub>alkyloxyC<sub>1</sub>-C<sub>6</sub>alkyl, NR<sub>11</sub>R<sub>12</sub> or C<sub>1</sub>-C<sub>6</sub>alkylamino-C<sub>1</sub>-C<sub>6</sub>alkyl, wherein R<sub>11</sub> and R<sub>12</sub> are independently selected from hydrogen, C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, arylcarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, C<sub>1</sub>-C<sub>6</sub>alkylcarboxy or arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy; wherein the alkyl and aryl groups are optionally substituted; or
- 10 R<sub>9</sub> and R<sub>10</sub> are independently a saturated or partial saturated cyclic 5, 6 or 7 membered amine, imide or lactam or a salt thereof with a pharmaceutically acceptable acid or base, or any optical isomer or mixture of optical isomers, including a racemic mixture, or any tautomeric forms.
- The compounds of Formula 1 are oxalylamide compounds having in common key structural features required of non hydrolysable protein tyrosine phosphatase inhibitors, most particularly PTP1B and/or TC-PTP inhibitors. These structural features endow the present compounds with the appropriate molecular shape necessary to fit into the enzymatic active site, to bind to such site in a non covalently way, thereby blocking the site and inhibiting enzymatic biological activity. Referring to Formula 1, such structural features include the oxalylamide and an ortho-carboxylic acid attached to a hydrophobic group, preferably an aryl as defined below The compounds of the invention can be further modified to act as prodrugs.

It is a well known problem in drug discovery that compounds, such as enzyme inhibitors, may be very potent and selective in biochemical assays, yet be inactive in vivo. This lack of so-called bioavailability may be ascribed to a number of different factors such as lack of or poor absorption in the gut, first pass metabolism in the liver, poor uptake in cells. Although the factors determining bioavailability are not completely understood, there are many examples in the scientific literature - well-known to those skilled in the art - of how to modify compounds, which are potent and selective in biochemical assays but show low or no activity in

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vivo, into drugs that are biologically active. It is within the scope of the invention to modify the compounds of the invention, termed the 'original compound' or "prototype", by attaching chemical groups that will improve the bioavailability of said compounds in such a way that the uptake in cells or mammals is facilitated. Examples of said modifications, which are not intended in any way to limit the scope of the invention, include changing of one or more carboxy groups to esters (for instance methyl esters, ethyl esters, acetoxymethyl esters or other acyloxymethyl esters). Compounds of the invention, original compounds, modified by attaching chemical groups are termed 'modified compounds' Said chemical groups may or may not be apparent in the claims of this invention. Other examples of modified compounds, which are not intended in any way to limit the scope of the invention, are compounds that have been cyclized at specific positions - so called 'cyclic compounds' - which upon uptake in cells or mammals become hydrolyzed at the same specific position(s) in the molecule to yield the compounds of the invention, the original compounds, which are then said to be 'non-cyclic' For the avoidance of doubt, it is understood that the latter original compounds in most cases will contain other cyclic or heterocyclic structures that will not be hydrolyzed after uptake in cells or mammals. Generally, said modified compounds will not show a behavior in biochemical assays similar to that of the original compound, i.e. the corresponding compounds of the invention without the attached chemical groups or said modifications. Said modified compounds may even be inactive in biochemical assays. However, after uptake in cells or mammals these attached chemical groups of the modified compounds may in turn be removed spontaneously or by endogenous enzymes or enzyme systems to yield compounds of the invention, original compounds. 'Uptake' is defined as any process that will lead to a substantial concentration of the compound inside cells or in mammals. After uptake in cells or mammals and after removal of said attached chemical group or hydrolysis of said cyclic compound, the compounds may have the same structure as the original compounds and thereby regain their activity and hence become active in cells and/or in vivo after uptake. A number of procedures, well known to those skilled in the art,

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may be used to verify that the attached chemical groups have been removed or that the cyclic compound has been hydrolyzed after uptake in cells or mammals. An example, which is not intended in any way to limit the scope of the invention, is given in the following. A mammalian cell line, which can be obtained from the American Tissue Type Collection or other similar governmental or commercial sources, is incubated with said modified compound. After incubation at conditions well known to those skilled in the art, the cells are washed appropriately, lysed and the lysate is isolated. Appropriate controls, well known to those skilled in the art, must be included. A number of different procedures, well known to those skilled in the art, may in turn be used to extract and purify said compound from said lysate. Said compound may or may not retain the attached chemical group or said cyclic compound may or may not have been hydrolyzed. Similarly, a number of different procedures - well known to those skilled in the art - may be used to characterize said purified compound structurally and chemically. Since said purified compound has been isolated from said cell lysate and hence has been taken up by said cell line, a comparison of said structurally and chemically characterized compound with that of the original unmodified compound (i.e. without said attached chemical group or said non-cyclic compound) will immediately provide to those skilled in the art information on whether the attached chemical group as been removed in the cell or whether the cyclic compound has been hydrolyzed. As a further analysis, said purified compound may be subjected to enzyme kinetic analysis as described in detail in the present invention. If the kinetic profile is similar to that of the original compound without said attached chemical group, but different from said modified compound, this confirms that said chemical group has been removed or said cyclic compounds has been hydrolyzed. Similar techniques may be used to analyze compounds of the invention in whole animals and mammals.

Preferred prodrug classes for the present compounds include acyloxymethyl esters or acyloxymethyl carbamates of the compounds of the present invention which may be prepared by the following general.

procedure (C.Schultz et al, J. Biol. Chem., 1993, 268, 6316-6322.) and (Alexander, J. et al, J. Med. Chem. 1991, 34, 78-81).

A carboxylic acid (1 equivalent) is suspended in dry acetonitrile (2 ml per 0.1 mmol). Diisopropyl amine (3.0 equivalents) is added followed by bromomethyl acetate (1.5 equivalents). The mixture is stirred under nitrogen overnight at room temperature. Acetonitrile is removed under reduced pressure to yield an oil which is diluted in ethyl acetate and washed with water (3 x). The organic layer is dried over anhydrous magnesium sulfate. Filtration followed by solvent removal under reduced pressure affords a crude oil. The product is purified by column chromatography on silica gel, using an appropriate solvent system.

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#### **DEFINITIONS**

As used herein, the term "attached" or "-" (e.g. –C(O)-R<sub>13</sub>, which indicates the carbonyl attachment point to the scaffold) signifies a stable covalent bond, certain preferred points of attachment points being apparent to those skilled in the art.

The terms "halogen" or "halo" include fluorine, chlorine, bromine, and iodine.

The term "alkyl" includes  $C_1$ - $C_6$  straight chain saturated, methylene and  $C_2$ - $C_6$  unsaturated aliphatic hydrocarbon groups,  $C_1$ - $C_6$  branched saturated and  $C_2$ - $C_6$  unsaturated aliphatic hydrocarbon groups,  $C_3$ - $C_6$  cyclic saturated and  $C_5$ - $C_6$  unsaturated aliphatic hydrocarbon groups, and  $C_1$ - $C_6$  straight chain or branched saturated and  $C_2$ - $C_6$  straight chain or branched unsaturated aliphatic hydrocarbon groups substituted with  $C_3$ - $C_6$  cyclic saturated and unsaturated aliphatic hydrocarbon groups having the specified number of carbon atoms. For example, this definition shall include but is not limited to methyl (Me), ethyl (Et), propyl (Pr), butyl (Bu), pentyl, hexyl, heptyl, ethenyl, propenyl, butenyl, penentyl, hexenyl,

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isopropyl (i-Pr), isobutyl (i-Bu), *tert*-butyl (*t*-Bu), *sec*-butyl (*s*-Bu), isopentyl, neopentyl, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclopentenyl, cyclohexenyl, methylcyclopropyl, ethylcyclohexenyl, butenylcyclopentyl,

cyclohexenyl, methylcyclopropyl, ethylcyclohexenyl, butenylcyclopentyl, and the like. The alkyl group as defined above is optionally substituted wherein the substitutents are independently selected from halo, cyano, nitro, trihalomethyl, carbamoyl, hydroxy, oxo, COOR3, CONR9R10, C1-C6alkyl, C1-C6alkyloxy, aryloxy, arylC1-C6alkyloxy, thio, C1-C6alkylthio, arylthio, arylC1-C6alkylthio, NR9R10, C1-C6alkylamino, arylamino, arylC1-C6alkylamino, di(arylC1-C6alkyl)amino, C1-C6alkylcarbonyl, arylC1-C6alkylcarboxy, arylcarboxy, arylC1-C6alkylcarboxy, C1-C6alkylcarbonylamino, -C1-C6alkylaminoCOR14, arylC1-C6alkylcarbonylamino, tetrahydrofuranyl, morpholinyl, piperazinyl, -CONR9R10, -C1-C6-alkylCONR9R10, or a saturated or partial saturated cyclic 5, 6 or 7 membered amine, imide or lactam; wherein R14 is hydroxy, C1-C6alkyl, aryl, arylC1-C6alkyl, C1-C6alkyloxy, aryloxy, arylC1-C6alkyloxy and R3 is defined as above or NR9R10, wherein R9, R10 are defined as above.

The term "saturated, partially saturated or aromatic cyclic, bicyclic or tricyclic ring system" represents but are not limit to aziridinyl, pyrrolyl, pyrrolinyl, pyrrolidinyl, imidazolyl, 2-imidazolinyl, imidazolidinyl, pyrazolyl, 2-pyrazolinyl, 1,2,3-triazolyl, 1,2,4-triazolyl, morpholinyl, piperidinyl, thiomorpholinyl, piperazinyl, indolyl, isoindolyl, 1,2,3,4-tetrahydro-quinolinyl, 1,2,3,4-tetrahydro-isoquinolinyl, 1,2,3,4-tetrahydro-quinoxalinyl, indolinyl, indazolyl, benzimidazolyl, benzotriazolyl, purinyl, carbazolyl, acridinyl, phenothiazinyl, phenoxazinyl, iminodibenzyl, iminostilbenyl.

The term "alkyloxy" (e.g. methoxy, ethoxy, propyloxy, allyloxy, cyclohexyloxy) represents an "alkyl" group as defined above having the indicated number of carbon atoms attached through an oxygen bridge. The term "alkyloxyalkyl" represents an "alkyloxy" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

The term "alkyloxyalkyloxy" represents an "alkyloxyalkyl" group attached through an oxygen atom as defined above having the indicated number of carbon atoms.

The term "aryloxy" (e.g. phenoxy, naphthyloxy and the like) represents an aryl group as defined below attached through an oxygen bridge.

The term "arylalkyloxy" (e.g. phenethyloxy, naphthylmethyloxy and the like) represents an "arylalkyl" group as defined below attached through an oxygen bridge.

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The term "arylalkyloxyalkyl" represents an "arylalkyloxy" group as defined above attached through an "alkyl" group defined above having the indicated number of carbon atoms.

The term "arylthio" (e.g. phenylthio, naphthylthio and the like) represents an "aryl" group as defined below attached through an sulfur bridge.

The term "alkyloxycarbonyl" (e.g. methylformiat, ethylformiat and the like) represents an "alkyloxy" group as defined above attached through a carbonyl group.

The term "aryloxycarbonyl" (e.g. phenylformiat, 2-thiazolylformiat and the like) represents an "aryloxy" group as defined above attached through a carbonyl group.

The term "arylalkyloxycarbonyl" (e.g. benzylformiat, phenyletylformiat and the like) represents an "arylalkyloxy" group as defined above attached through a carbonyl group.

The term "alkyloxycarbonylalkyl" represents an "alkyloxycarbonyl" group as defined above attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

The term "arylalkyloxycarbonylalkyl" represents an "arylalkyloxycarbonyl" group as defined above attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

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The term "alkylthio" (e.g. methylthio, ethylthio, propylthio, cyclohexenylthio and the like) represents an "alkyl" group as defined above having the indicated number of carbon atoms attached through a sulfur bridge.

The term "arylalkylthio" (e.g. phenylmethylthio, phenylethylthio, and the like) represents an "arylalkyl" group as defined above having the indicated number of carbon atoms attached through a sulfur bridge.

The term "alkylthioalkyl" represents an "alkylthio" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

10 The term "arylalkylthioalkyl" represents an "arylalkylthio" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

The term "alkylamino" (e.g. methylamino, diethylamino, butylamino, Npropyl-N-hexylamino, (2-cyclopentyl)propylamino, hexenylamino, pyrrolidinyl, piperidinyl and the like) represents one or two "alkyl" groups as defined above having the indicated number of carbon atoms attached through an amine bridge. The two alkyl groups may be taken together with the nitrogen to which they are attached forming a saturated, partially saturated or aromatic cyclic, bicyclic or tricyclic ring system containing 3 to 14 carbon atoms and 0 to 3 additional heteroatoms selected from nitrogen, oxygen or sulfur, the ring system can optionally be substituted with at least one C<sub>1</sub>-C<sub>6</sub>alkyl, aryl, arylC<sub>1</sub>-C<sub>6</sub>alkyl, hydroxy, oxo, C<sub>1</sub>-C<sub>6</sub>alkyloxy, C<sub>1</sub>-C<sub>6</sub>alkyloxyC<sub>1</sub>-C<sub>6</sub>alkyl, NR<sub>9</sub>R<sub>10</sub>, C<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl substituent wherein the alkyl and aryl groups are optionally substituted as defined in the definition section and  $R_9$  and  $R_{10}$  are defined as above. The term "arylalkylamino" (e.g. benzylamino, diphenylethylamino and the like) represents one or two "arylalkyl" groups as defined above having the indicated number of carbon atoms attached through an amine bridge. The two "arylalkyl" groups may be taken together with the nitrogen to which they are attached forming a saturated, partially saturated or aromatic cyclic, bicyclic or tricyclic ring system containing 3 to 14 carbon atoms and 0 to 3 additional heteroatoms selected from nitrogen, oxygen or sulfur, the ring system can optionally be substituted with at least one C<sub>1</sub>-C<sub>6</sub>alkyl, aryl,

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arylC<sub>1</sub>-C<sub>6</sub>alkyl, hydroxy, oxo, C<sub>1</sub>-C<sub>6</sub>alkyloxy, C<sub>1</sub>-C<sub>6</sub>alkyloxyC<sub>1</sub>-C<sub>6</sub>alkyl, NR<sub>9</sub>R<sub>10</sub>, C<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl substituent wherein the alkyl and aryl groups are optionally substituted as defined in the definition section and R<sub>9</sub> and R<sub>10</sub> are defined as above.

The term "alkylaminoalkyl" represents an "alkylamino" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

The term "arylalkylaminoalkyl" represents an "arylalkylamino" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

The term "arylalkyl" (e.g. benzyl, phenylethyl) represents an "aryl" group as defined below attached through an alkyl having the indicated number of carbon atoms or substituted alkyl group as defined above.

The term "alkylcarbonyl" (e.g. cyclooctylcarbonyl, pentylcarbonyl, 3-

hexenylcarbonyl) represents an "alkyl" group as defined above having the indicated number of carbon atoms attached through a carbonyl group.

The term "arylcarbonyl" (benzoyl) represents an "aryl" group as defined above attached through a carbonyl group.

The term "arylalkylcarbonyl" (e.g. phenylcyclopropylcarbonyl, phenylethylcarbonyl and the like) represents an "arylalkyl" group as

defined above having the indicated number of carbon atoms attached through a carbonyl group.

The term "alkylcarbonylalkyl" represents an "alkylcarbonyl" group attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

The term "arylalkylcarbonylalkyl" represents an "arylalkylcarbonyl" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

The term "alkylcarboxy" (e.g. heptylcarboxy, cyclopropylcarboxy, 3-pentenylcarboxy) represents an "alkylcarbonyl" group as defined above wherein the carbonyl is in turn attached through an oxygen bridge.

The term "arylcarboxyalkyl" (e.g. phenylcarboxymethyl) represents an

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"arylcarbonyl" group defined above wherein the carbonyl is in turn attached through an oxygen bridge to an alkyl chain having the indicated number of carbon atoms.

The term "arylalkylcarboxy" (e.g. benzylcarboxy, phenylcyclopropylcarboxy and the like) represents an "arylalkylcarbonyl" group as defined above wherein the carbonyl is in turn attached through an oxygen bridge.

The term "alkylcarboxyalkyl" represents an "alkylcarboxy" group attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

The term "arylalkylcarboxyalkyl" represents an "arylalkylcarboxy" group attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

The term "alkylcarbonylamino" (e.g. hexylcarbonylamino,

cyclopentylcarbonyl-aminomethyl, methylcarbonylaminophenyl) represents an "alkylcarbonyl" group as defined above wherein the carbonyl is in turn attached through the nitrogen atom of an amino group. The nitrogen atom may itself be substituted with an alkyl or aryl group.

The term "arylalkylcarbonylamino" (e.g. benzylcarbonylamino and the like) represents an "arylalkylcarbonyl" group as defined above wherein the carbonyl is in turn attached through the nitrogen atom of an amino group. The nitrogen atom may itself be substituted with an alkyl or aryl group. The term "alkylcarbonylaminoalkyl" represents an "alkylcarbonylamino" group attached through an "alkyl" group as defined above having the indicated number of carbon atoms. The nitrogen atom may itself be substituted with an alkyl or aryl group.

The term "arylalkylcarbonylaminoalkyl" represents an "arylalkylcarbonylamino" group attached through an "alkyl" group as defined above having the indicated number of carbon atoms. The nitrogen atom may itself be substituted with an alkyl or aryl group.

The term "alkylcarbonylaminoalkylcarbonyl" represents an alkylcarbonylaminoalkyl group attached through a carbonyl group. The nitrogen atom may be further substituted with an "alkyl" or "aryl" group.

The term "aryl" represents a substituted or unsubstituted, mono-, di- or trisubstituted monocyclic, polycyclic, biaryl and heterocyclic aromatic groups covalently attached at any ring position capable of forming a stable covalent bond, certain preferred points of attachment being apparent to those skilled in the art (e.g., 3-indolyl, 4-imidazolyl). The aryl substituents are independently selected from the group consisting of halo, nitro, cyano, trihalo-methyl,  $C_1$ - $C_6$ alkyl, aryl, aryl $C_1$ - $C_6$ alkyl, hydroxy, COOR<sub>3</sub>, CONR<sub>9</sub>R<sub>10</sub>,  $C_1$ - $C_6$ alkyloxy,  $C_1$ - $C_6$ alkyloxy $C_1$ - $C_6$ alkyloxy, aryl $C_1$ - $C_6$ alkyloxy

C<sub>6</sub>alkyloxy, arylC<sub>1</sub>-C<sub>6</sub>alkyloxyC<sub>1</sub>-C<sub>6</sub>alkyl, thio, C<sub>1</sub>-C<sub>6</sub>alkylthio, C<sub>1</sub>
C<sub>6</sub>alkylthioC<sub>1</sub>-C<sub>6</sub>alkyl, arylthio, arylC<sub>1</sub>-C<sub>6</sub>alkylthio, arylC<sub>1</sub>-C<sub>6</sub>alkylthioC<sub>1</sub>
C<sub>6</sub>alkyl, NR<sub>9</sub>R<sub>10</sub>, C<sub>1</sub>-C<sub>6</sub>-alkylamino, C<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, arylamino, arylC<sub>1</sub>-C<sub>6</sub>alkylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, di(arylC<sub>1</sub>-

C<sub>6</sub>alkyl)aminoC<sub>1</sub>-C<sub>6</sub>alkyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, C<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylC<sub>1</sub>-C<sub></sub>

15 C<sub>6</sub>alkylcarboxy, C<sub>1</sub>-C<sub>6</sub>alkylcarboxy-C<sub>1</sub>-C<sub>6</sub>alkyl, arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxy, arylC<sub>1</sub>-C<sub>6</sub>alkylcarboxyC<sub>1</sub>-C<sub>6</sub>alkyl, carboxyC<sub>1</sub>-C<sub>6</sub>alkyl-oxy, C<sub>1</sub>-C<sub>6</sub>alkylcarbonylamino, C<sub>1</sub>-C<sub>6</sub>alkylcarbonylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, -carbonylNR<sub>7</sub>C<sub>1</sub>-C<sub>6</sub>alkylCOR<sub>14</sub>, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylamino, arylC<sub>1</sub>-C<sub>6</sub>alkylcarbonylaminoC<sub>1</sub>-C<sub>6</sub>alkyl, -CONR<sub>9</sub>R<sub>10</sub>, or -C<sub>1</sub>-C<sub>6</sub>alkylCONR<sub>9</sub>R<sub>10</sub>.

wherein R<sub>3</sub>, R<sub>9</sub>, R<sub>10</sub>, and R<sub>14</sub> are defined as above and the alkyl and aryl groups contained therein are optionally substituted as defined above. The definition of aryl includes but is not limited to phenyl, biphenyl, indenyl, fluorenyl, naphthyl (1-naphthyl, 2-naphthyl), pyrrolyl (2-pyrrolyl), pyrazolyl (3-pyrazolyl), imidazolyl (1-imidazolyl, 2-imidazolyl, 4-imidazolyl,

5-imidazolyl), triazolyl (1,2,3-triazol-1-yl, 1,2,3-triazol-2-yl 1,2,3-triazol-4-yl, 1,2,4-triazol-3-yl), oxazolyl (2-oxazolyl, 4-oxazolyl, 5-oxazolyl), isoxazolyl (3-isoxazolyl, 4-isoxazolyl, 5-isoxazolyl), thiazolyl (2-thiazolyl, 4-thiazolyl, 5-thiazolyl), thiophenyl (2-thiophenyl, 3-thiophenyl, 4-thiophenyl, 5-thiophenyl), furanyl (2-furanyl, 3-furanyl, 4-furanyl, 5-furanyl), pyridyl (2-

pyridyl, 3-pyridyl, 4-pyridyl, 5-pyridyl), 5-tetrazolyl, pyrimidinyl (2-pyrimidinyl, 4-pyrimidinyl, 5-pyrimidinyl, 6-pyrimidinyl), pyrazinyl, pyridazinyl (3-pyridazinyl, 4-pyridazinyl, 5-pyridazinyl), quinolyl (2-quinolyl, 3-quinolyl, 4-quinolyl, 5-quinolyl, 7-quinolyl, 8-quinolyl), isoquinolyl (1-isoquinolyl, 3-isoquinolyl, 4-isoquinolyl, 5-isoquinolyl, 6-

isoquinolyl, 7-isoquinolyl, 8-isoquinolyl), benzo[b]füranyl (2benzo[b]furanyl, 3-benzo[b]furanyl, 4-benzo[b]furanyl, 5-benzo[b]furanyl, 6-benzo[b]furanyl, 7-benzo[b]furanyl), 2,3-dihydro-benzo[b]furanyl (2-(2,3dihydro-benzo[b]furanyl), 3-(2,3-dihydro-benzo[b]furanyl), 4-(2,3-dihydrobenzo[b]furanyl), 5-(2,3-dihydro-benzo-[b]furanyl), 6-(2,3-dihydro-benzo-[b]furanyl), 7-(2,3-dihydro-benzo[b]furanyl)), benzo[b]thiophenyl (2benzo[b]thiophenyl, 3-benzo[b]thiophenyl, 4-benzo[b]thiophenyl, 5benzo[b]thiophenyl, 6-benzo[b]thiophenyl, 7-benzo[b]thiophenyl), 2,3dihydro-benzo[b]-thiophenyl (2-(2,3-dihydro-benzo[b]thiophenyl), 3-(2,3dihydro-benzo[b]-thiophenyl), 4-(2,3-dihydro-benzo[b]thiophenyl), 5-(2,3-10 dihydro-benzo[b]-thiophenyl), 6-(2,3-dihydro-benzo[b]thiophenyl), 7-(2,3dihydro-benzo[b]-thiophenyl)), 4,5,6,7-tetrahydro-benzo[b]thiophenyl (2-(4,5,6,7-tetrahydro-benzo-[b]thiophenyl), 3-(4,5,6,7-tetrahydro-benzo-[b]thiophenyl), 4-(4,5,6,7-tetrahydro-benzo[b]thiophenyl), 5-(4,5,6,7tetrahydro-benzo-[b]thiophenyl), 6-(4,5,6,7-tetrahydro-benzo-15. [b]thiophenyl), 7-(4,5,6,7-tetrahydro-benzo[b]thiophenyl)), 4,5,6,7tetrahydro-thieno[2,3-c]pyridyl (4-(4,5,6,7-tetrahydro-thieno[2,3-c]pyridyl), 5-4,5,6,7-tetrahydro-thieno[2,3-c]pyridyl), 6-(4,5,6,7-tetrahydro-thieno[2,3c]pyridyl), 7-(4,5,6,7-tetrahydro-thieno[2,3-c]pyridyl)), indolyl (1-indolyl, 2indolyl, 3-indolyl, 4-indolyl, 5-indolyl, 6-indolyl, 7-indolyl), isoindolyl (1-20 isoindolyl, 2-isoindolyl, 3-isoindolyl, 4-isoindolyl, 5-isoindolyl, 6-isoindolyl, 7-isoindolyl), 1,3-dihydro-isoindolyl (1-(1,3-dihydro-isoindolyl), 2-(1,3dihydro-isoindolyl), 3-(1,3-dihydro-isoindolyl), 4-(1,3-dihydro-isoindolyl), 5-(1,3-dihydro-isoindolyl), 6-(1,3-dihydro-isoindolyl), 7-(1,3-dihydroisoindolyl)), indazole (1-indazolyl, 3-indazolyl, 4-indazolyl, 5-indazolyl, 6indazolyl, 7-indazolyl), benzimidazolyl (1-benzimidazolyl, 2-benzimidazolyl, 4-benzimidazolyl, 5-benzimidazolyl, 6-benzimidazolyl, 7-benzimidazolyl, 8benzimidazolyl), benzoxazolyl (1-benz-oxazolyl, 2-benzoxazolyl), benzothiazolyl (1-benzothiazolyl, 2-benzo-thiazolyl, 4-benzothiazolyl, 5-30 benzothiazolyl, 6-benzothiazolyl, 7-benzothiazolyl), carbazolyl (1carbazolyl, 2-carbazolyl, 3-carbazolyl, 4-carbazolyl), 5H-dibenz[b,f]azepine (5H-dibenz[b,f]azepin-1-yl, 5H-dibenz-[b,f]azepine-2-yl, 5Hdibenz[b,f]azepine-3-yl, 5H-dibenz-[b,f]azepine-4-yl, 5H-dibenz[b,f]azepine-5-yl), 10,11-dihydro-5H-dibenz[b,f]azepine (10,11-dihydro-5H-

dibenz[b,f]azepine-1-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-2-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-3-yl, 10,11-dihydro-5H-dibenz-[b,f]azepine-4-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-5-yl), piperidinyl (2-piperidinyl, 3-piperidinyl, 4-piperidinyl), pyrrolidinyl (1-pyrrolidinyl, 2-pyrrolidinyl, 3-pyrrolidinyl), phenylpyridyl (2-phenyl-pyridyl, 3-phenyl-pyridyl, 4-phenylpyridyl), phenylpyrimidinyl (2-phenylpyrimidinyl, 4-phenyl-pyrimidinyl, 5-phenylpyrimidinyl, 6-phenylpyrimidinyl), phenylpyrazinyl, phenylpyridazinyl (3-phenylpyridazinyl, 4-phenylpyridazinyl, 5-phenyl-pyridazinyl).

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The term "arylcarbonyl" (e.g. 2-thiophenylcarbonyl, 3-methoxy-anthrylcarbonyl, oxazolylcarbonyl) represents an "aryl" group as defined above attached through a carbonyl group.

The term "arylalkylcarbonyl" (e.g. (2,3-dimethoxyphenyl)propylcarbonyl, (2-chloronaphthyl)pentenylcarbonyl, imidazolylcyclopentylcarbonyl) represents an "arylalkyl" group as defined above wherein the "alkyl" group is in turn attached through a carbonyl.

The term "aryloxyalkyl" represents an "aryloxy" group as defined above attached through an "alkyl" group defined above having the indicated number of carbon atoms.

The term "arylaminocarbonylaminoalkyl" represents an "arylaminocarbonylamino" group as defined above attached through an "alkyl" group as defined above having the indicated number of carbon atoms.

The term "R8R9Nalkyl" is as defined under "substituted alkyl" or "optionally substituted alkyl".

The term "arylaminoalkyl" represents an "arylamino" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

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The term "arylcarbonylaminoalkyl" represents an "arylcarbonylamino" group attached through an alkyl group as defined above having the indicated number of carbon atoms.

As used herein, the term "vicinity" applied with respect to the active site of a PTPase means the space occupied by a half sphere – with its apex pointing towards aspartic acid 48 - having its center in the side chain nitrogen atom of the guanidinium group of residue 221 (arginine), which points away from the phosphate binding loop (residue Arg221 to Cys215). The radius of the half sphere is 27 Å.

As used herein, the term "structurally similar" means any PTPase that contains an aspartic acid in residue position 48 (PTP1B numbering – as defined in Chernoff et al, 1989, *supra*) and is more than 50 % identical and preferably more than 65 % identical and most preferably more than 80 % identical to PTP1B (Chernoff et al., *supra*) and/or TC-PTP (Cool et al., Proc. Natl. Acad. Sci. U.S.A. 86: 5257-5261 (1989)) at the primary amino acid sequence level in the catalytic domain as defined below. Percent indentity can be determined using standard algorithms e.g. BLAST, BLASTP MEGALIGN, etc using default parameters.

As used herein, the term "catalytic domain" means the primary amino acid sequence of a PTPase that corresponds to the primary amino acid sequence between Asn 40 and Gln 262 (both residues included) in PTP1B (Chernoff et al., *supra*).

As used herein, the term "centroid" means the position for the stated atoms calculated by averaging the x coordinates of the atoms to obtain the x coordinate of the centroid, averaging the y coordinates of the atoms to obtain the y coordinate of the centroid, and averaging the z coordinates of the atoms to obtain the z coordinate of the centroid.

As used herein, the term "phosphate isostere" means a chemical group, which binds to one or more of the side chains or the main chain of the residues in the so-called P-loop or PTP signature motif of PTPases (i.e. Cys215-Xxx216-Xxx217-Xxx218-Xxx219-Xxx220-Arg221, where Cys215 and Arg221 are absolutely conserved, whereas Xxx stands for less conserved residues). In PTP1B the P-loop residues are: Cys215-

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Ser216-Ala217-Gly218-Ile219-Gly220-Arg221). As a non limiting example the following groups are phosphate isosteres: -CH<sub>2</sub>PO(OH)<sub>2</sub>, -CHFPO(OH)<sub>2</sub>, -CF<sub>2</sub>PO(OH)<sub>2</sub>, -NHCOCOOH, -OCH(COOH)<sub>2</sub>, -OCF(COOH)<sub>2</sub>, -OCH<sub>2</sub>COOH. -CONHCH<sub>2</sub>COOH, -CONHCHFCOOH and -CONHCF<sub>2</sub>COOH.

As used herein, the term "carboxylic acid isostere" means a compound resembling a carboxy group in its electronic and steric configuration and in its biological action (effecting inhibition of the class of structurally similar PTPases) but having a different chemical structure. As a non limiting example, the following residues and heterocycles are carboxylic acid isosteres: -CONH<sub>2</sub>, -SONH<sub>2</sub>, -SO<sub>2</sub>NH<sub>2</sub>.

As used herein the term "interact" or "interaction" when used in the context of a moiety or group of an inhibitor interacting with the active site or vicinity thereof of a PTPase, means the formation of noncovalent bonds, such as hydrogen bonds, salt bridges, hydrophobic interactions van der Waals forces, cation  $\pi$  interactions, or  $\pi$ ,  $\pi$  interactions, aromatic-aromatic interactions, (Copeland, Enzymes-a practical introduction to structure, mechanism, and data analysis, VCH Publishers, Inc..New York (1996)) or by forming covalent bonds. Preferably, interactions between inhibitors of the invention and PTPs occur through non-covalent bonds.

As used herein, the term "hydrophobic" means a nonpolar chemical group (e.g. phenyl, naphthyl, cyclopropyl, cyclobutyl, cyclobacyl, tert-butyl, isopropyl as nonlimiting examples) when present in the aqueous

phase, in the vicinity of an enzyme, its hydrocarbon framework disturbs the degree of randomness of the water molecules, which forces the water molecules to associate by hydrogen bonding to form quasi-crystalline clusters or "ice-bergs". This localized increase in the ordered structure of water will result in a loss of entropy, accompanied by an increase in the free energy of the system. Thus, a driving force operates to reject the hydrocarbon region of the drug/inhibitor from the aqueous phase so that binding to one or more similar hydrocarbon chain(s) within the enzyme molecule is facilitated.

As used herein, the term "hydrogen bond" means an association between an electronegative atom, e.g. fluorine, oxygen, nitrogen, or sulfur, and a hydrogen atom attached to another such electronegative atom.

As used herein, the term "salt bridge" means any electrostatic bond between positively and negatively charged groups.

The compounds of the present invention have asymmetric centers and may occur as racemates, racemic mixtures, and as individual enantiomers or diastereoisomers, with all isomeric forms being included in the present invention as well as mixtures thereof.

Pharmaceutically acceptable salts of the compounds of formula 1, where a basic or acidic group is present in the structure, are also included within the scope of this invention. When an acidic substituent is present, such as -COOH, 5-tetrazolyl or -P(O)(OH)<sub>2</sub>, there can be formed the ammonium, morpholinium, sodium, potassium, barium, calcium salt, and the like, for use as the dosage form. When a basic group is present, such as amino or a basic heteroaryl radical, such as pyridyl, an acidic salt, such as hydrochloride, hydrobromide, phosphate, sulfate, trifluoroacetate, trichloroacetate, acetate, oxalate, maleate, pyruvate, malonate, succinate, citrate, tartarate, fumarate, mandelate, benzoate, cinnamate, methanesulfonate, ethane sulfonate, picrate and the like, and include acids related to the pharmaceutically acceptable salts listed in Journal of Pharmaceutical Science, 66, 2 (1977) and incorporated herein by reference, can be used as the dosage form.

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Also, in the case of the -COOH or -P(O)(OH)<sub>2</sub> being present, pharmaceutically acceptable esters can be employed, e.g., methyl, *tert*-butyl, acetoxymethyl, pivaloyloxymethyl, and the like, and those esters known in the art for modifying solubility or hydrolysis characteristics for use as sustained release or prodrug formulations.

In addition, some of the compounds of the instant invention may form solvates with water or common organic solvents. Such solvates are encompassed within the scope of the invention.

As used herein, "treatment" shall include therapeutic or preventative management, treatment, cure, or palliation of a disease state or a measurable delay in its onset or recurrence or measurable reduction in its severity.

The term "therapeutically effective amount" shall mean that amount of drug or pharmaceutical agent that will elicit the biological or medical response of a tissue, system, animal, or human that is being sought by a researcher, veterinarian, medical doctor or other other biological or clinical investigator. Also included in the present invention is a process for isolation of PTPases via affinity purification procedures based on the use of immobilized compounds of the invention. Isolation can be effected using procedures otherwise well-known to those skilled in the art. Such methods, may be used to identify novel PTPases or other molecules with phosphotyrosine recognition units and to elucidate the function of both novel and previously identified PTPases. As a nonlimiting example, compounds of the invention may be immobilized by coupling to a solid-phase support, such as as exemplified in examples 119 and 120. See also Example 121. A tissue sample or a sample from a cell line prepared as a lysate by methods well-known to those skilled in the art may be passed over said solid-phase coupled with a compound of the invention. After appropriate washing procedures designed to remove material that binds nonspecifically to said solidphase, using standard procedures well known to those skilled in the art, mostly-PTPases or other molecules with phosphotyrosine recognition units will be bound to the compounds of the invention coupled to the

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solid phase. Said PTPases or other molecules with phosphotyrosine recognition units may in turn be released by procedures well-known in the art and further subjected to amino acid sequence analysis according to standard procedures well-known to those skilled in the art.

5 By back-translation of said amino acid sequence into a nucleotide sequence of the corresponding cDNA can be deduced using the appropriate genetic code. Said nucleotide sequence can be used to design and produce an equivalent oligonucleotide, which in turn can be used to identify partial or full-length cDNA clones from appropriate cDNA libraries encoding a protein or glycoprotein corresponding to or

cDNA libraries encoding a protein or glycoprotein corresponding to or similar to the isolated PTPase or molecule with pTyr recognition units. Said oligonucleotide or isolated cDNA clone(s) can similarly be used to isolate genomic clones corresponding to said cDNA clones. Said partial or full-length cDNA can be inserted into appropriate vectors and expressed and purified proteins with procedures well known to those skilled in the art. Said purified proteins, in particular PTPases, may be

The invention is further directed to compounds of the invention coupled to a suitable solid-phase matrix such as a Wang-resin or a Rink-resin, e.g., for further synthesis, combinational synthesis, or as a support for affinity purification.

used to further analyze the inhibitory capacity and selectivity of

compounds of the invention as described.

The invention is further directed to a method for isolating a protein or a glycoprotein with affinity for a compound according to the invention from a biological sample, comprising:

- contacting a compound of the invention immobilized by coupling to a suitable solid-phase matrix with said biological sample in order for said immobilized compound to form a complex by binding said protein or glycoprotein,
- removing unbound material from said biological sample and isolating said complex, and
  - extracting said protein or glycoprotein from said complex.

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The invention is further directed to a method for isolating a protein-tyrosine phosphatase with affinity for a compound according to the invention from a biological sample, comprising

- contacting a compound of the invention immobilized by coupling to a suitable solid-phase matrix with said biological sample in order for said immobilized compound to form a complex by binding said protein-tyrosine phosphatase
  - removing unbound material from said biological sample and isolating said complex
- extracting said protein-tyrosine phosphatase.

The following compounds are encompassed by the invention: 5-(4-Chloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 7-(2,4-Dioxo-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 5-(4,5,6,7-Tetrachloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-isoindol-2-ylmethyl

- amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 5-(1,3-Dioxo-1,3-dihydro-benzo[f]isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- Oxalic acid (3-carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl) ester methyl ester;
- Oxalic acid (3-carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl) ester;
  - 7-Hydroxymethyl-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 7-(((Benzo[1,3]dioxole-5-carbonyl)-amino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  5-(3-Imidazol-1-yl-2,5-dioxo-pyrrolidin-1-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;

- 2-(Oxalyl-amino)-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 2-(Oxalyl-amino)-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5 2-(Oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 7-ethyl ester;
  - 7-Benzylcarbamoyl-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-2-(oxalyl-
- amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  5-(4-(4-Chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 7-(1,3-Dioxo-1,3-dihydro-isoindol-2-yloxymethyl)-2-(oxalyl-amino)-4,7-
- dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 7-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 7-(3-(2,4-Dimethoxy-phenyl)-ureidomethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 2-((3-Carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl)-carbamoyl)-nicotinic acid;
  - 5-(4-Fluoro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 25 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-
  - amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 30 7-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 7-(5,7-Dioxo-5,7-dihydro-[1,3]dioxolo[4,5-f]isoindol-6-ylmethyl2-(oxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;

- 7-(2,4-Dioxo-5-pyridin-2-ylmethylene-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 7-(2,4-Dioxo-5-pyridin-2-ylmethyl-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 7-(5-(4-Methoxy-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-(5-(4-Acetylamino-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-(5-(3,5-Dimethoxy-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl)-2-
- 10 (oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 7-(5-(1H-Imidazol-4(5)-ylmethylene)-2,4-dioxo-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid; 7-((2-(4-Methanesulfonyl-phenyl)-acetylamino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5-(1,3-Dioxo-4,7-epoxido-1,3,4,5,6,7-hexahydro-isoindol-2-ylmethyl)-2(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-((2-Amino-3-phenyl-propionylamino)methyl)-2-(oxalyl-amino)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-(((2R)-2-Amino-3-phenyl-propionylamino)-methyl)-2-(oxalyl-amino)-4,7-
- dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-((2-Acetylamino-3-(4-hydroxy-phenyl)-propionylamino)-methyl)-2-(oxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  7-((2-Acetylamino-3-methyl-butyrylamino)methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5-(5-Acetylamino-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  5-(4-Acetylamino-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-2-(oxalyl-amino)-
- 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
   5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-c]pyridin-6-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
   5-(5-Nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;

- 5-(5-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5-(4-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5 5-(4-Nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 2-(Oxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 2-(Oxalyl-amino)-7-(3-oxo-3H-benzo[d]isoxazol-2-ylmethyl)-4,7-dihydro-
- 10 5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 6-ethyl ester;
  - 5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 15 (L)-5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - .5-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 20 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 2-(Oxalyl-amino)-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 25 7-(((Benzo[1,3]dioxole-5-carbonyl)amino)methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 5-(4-(4-Chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 30 7-(3-(2,4-Dimethoxy-phenyl)-ureidomethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
  - 7-((2-(4-Methanesulfonyl-phenyl)acetylamino)methyl)-2-(oxalyl-amino)-
  - 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;

- 7-((2-Acetylamino-3-(4-hydroxy-phenyl)propionylamino)methyl)-2-(oxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid;
- 5-(S)-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 7-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 2-(Oxalyl-amino)-5-(S)-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  5-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-methyl-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  5-((1,1-Dioxo-1H-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino)-
- 7-((1,1-Dioxo-1H-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

- 5-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 20 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(7-Benzyloxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-
  - 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 25 5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-
  - (oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-
- 30 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-
  - (oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

- 6-(4-Methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1H-indol-3-yl)-acetylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 5-(R)-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
- 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
   5-(S)-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(S)-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-
  - 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 2-(S)-(Oxalyl-amino)-5-((4-phenoxy-benzylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 5-(S)-((4-Acetylamino-benzylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(S)-((Acetyl-(4-phenoxy-benzyl)amino)methyl)-2-(oxalyl-amino)-4,5,6,7-
- tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 7-(S)-((Acetyl-benzyl-amino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid;
    - 5-(S)-((1,1-Dioxo-1H-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid; 5-(6-Methoxy-4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid; 2-(Oxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-
- ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3-carboxylic acid;
  2-(Oxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3-carboxylic acid;
  7 (R) Carbamoyl 2 (oxalyl-amino)-4 5 6 7-tetrahydro-thieno[2,3-c]pyridin
  - 7-(R)-Carbamoyl-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
- 30 2-(Oxalyl-amino)-5-(S)-(2-oxo-tetrahydro-thiophen-3-ylcarbamoyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;
  - 2-(Oxalyl-amino)-5-(S)-phenylcarbamoyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

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2-(Oxalyl-amino)-7-(R)-phenylcarbamoyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

5-(R),7-(R)-Bis-benzyloxymethyl-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid;

6-Benzyl-2-(oxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1,6-benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid; or a pharmaceutically acceptable salt thereof

#### PHARMACOLOGICAL METHODS

The compounds are evaluated for biological activity with a truncated form of PTP1B (corresponding to the first 321 amino acids), which was expressed in E. coli and purified to apparent homogeneity using published procedures well-known to those skilled in the art. The enzyme reactions are carried out using standard conditions essentially as described by Burke et al. (Biochemistry 35; 15989-15996 (1996)) incorporated by reference. The assay conditions are as follows. Appropriate concentrations of the compounds of the invention (e.g., 0.1 to 100µM) are added to the reaction mixtures containing different concentrations of the substrate, p-nitrophenyl phosphate (range: 0.16 to 10 mM - final assay concentration). The buffer used was 50 mM HEPES pH 7.0, 100 mM sodium chloride, 0.1 % (w/v) bovine serum albumin, 5 mM glutathione, and 1 mM EDTA. The reaction was started by addition of the enzyme and carried out in microtiter plates at 25° C for 60 minutes. The reactions are stopped by addition of NaOH. The enzyme activity was determined by measurement of the absorbance at 405 nm with appropriate corrections for absorbance at 405 nm of the compounds and p-nitrophenyl phosphate. The data are analyzed using nonlinear regression fit to classical Michaelis Menten enzyme kinetic models. Inhibition is expressed as K<sub>i</sub> values in µM. The results of representative experiments are shown in Table 6.

#### Tabl 6

Inhibition of classical PTPases by compounds



K<sub>i</sub> (μM) at pH 7

* *	PTP1B	TC-PTP	ΡΤΡ α	ΡΤΡ β	PTP ε
	residue 48	residue 48	residue 48	residue 48	residue 48
Example No.	Asp	Asp	Asn	Asn	Asn
48	0.25		900	47	380
49	0.085			8.6	
50	0.07		1000	8 .	7- ×
52	1.2	•	> 400	107	> 500

# THE SYNTHESIS OF THE COMPOUNDS

In accordance with one aspect of the invention, compounds of the invention are prepared as illustrated in the following reaction schemes wherein n, m, X, Y, R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub> and R<sub>6</sub> are defined as above:

## 10 Method A

- a) NCCH<sub>2</sub>COOR<sub>3</sub>, sulphur, morpholine or triethylamine, ethanol; b)
- 15 R<sub>3</sub>OCOCOimidazole, tetrahydrofuran; c) 25 % trifluoroacetic acid/dichloromethane.

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The pharmaceutical carrier employed may be a conventional solid or liquid carrier. Examples of solid carriers are lactose, terra alba, sucrose, talc, gelatine, agar, pectin, acacia, magnesium stearate and stearic acid. Examples of liquid carriers are syrup, peanut oil, olive oil, water, and physiologic saline.

Similarly, the carrier or diluent may include any material that impacts controlled release of taste-masking properties, known to the art, such as glyceryl monostearate or glyceryl distearate, alone or mixed with a wax.

If a solid carrier for oral administration is used, the preparation can be tabletted, placed in a hard gelatine capsule in powder or pellet form or it can be in the form of a troche or lozenge. The amount of solid carrier will vary widely but will usually be from about 25 mg to about 1 g. If a liquid carrier is used, the preparation may be in the form of a syrup, emulsion, soft gelatin capsule or sterile injectable liquid such as an aqueous or non-aqueous liquid suspension or solution.

20 Generally, the compounds of this invention are dispensed in unit dosage form comprising 10-200 mg of active ingredient in or together with a pharmaceutically acceptable carrier per unit dosage.

The dosage of the compounds according to this invention is 1-500 mg/day, e.g. about 100 mg per dose, when administered to patients, e.g. humans, as a drug.

A typical tablet that may be prepared by conventional tabletting techniques contains

#### 30 <u>Core:</u>

Active compound (as free compound	100 mg	
or salt thereof)		
Colloidal silicon dioxide (Areosil®)	1.5 mg	
Cellulose, microcryst. (Avicel®)	70 mg	

Modified cellulose gum (Ac-Di-Sol®)

Magnesium stearate

7.5 mg

#### Coating:

5 HPMC

approx.

9 mg

\*Mywacett® 9-40 T

0.9 mg

\*Acylated monoglyceride used as plasticiser for film coating.

The route of administration may be any route which effectively transports the active compound to the appropriate or desired site of action, such as oral or parenteral e.g. rectal, transdermal, subcutaneous, intranasal, intramuscular, topical, intravenous, intraurethral, ophthalmic solution or an ointment, the oral route being preferred.

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#### **EXAMPLES**

The process for preparing compounds of Formula 1 and preparations containing them is further illustrated in the following examples, which, however, are not to be construed as limiting.

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experimental section.

Hereinafter, TLC is thin layer chromatography, CDCl $_3$  is deuterio chloroform, CD $_3$ OD is tetradeuterio methanol and DMSO-d $_6$  is hexadeuterio dimethylsulfoxide. The structures of the compounds are confirmed by either elemental analysis or NMR, where peaks assigned to characteristic protons in the title compounds are presented where appropriate.  $^1$ H NMR shifts ( $\delta_H$ ) are given in parts per million (ppm) down field from tetramethylsilane as internal reference standard. M.p.: is melting point and is given in  $^{\circ}$ C and is not corrected. Column chromatography was carried out using the technique described by W.C. Still *et al.*, *J. Org. Chem. 43:* 2923 (1978) on Merck silica gel 60 (Art. 9385). HPLC analyses are performed using  $5\mu m$  C18 4 x 250 mm column eluted with various mixtures of water and acetonitrile, flow = 1 ml/min, as described in the

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Wang-resin is polystyrene with a 4-hydroxymethylphenol ether linker. Compounds used as starting material are either known compounds or compounds which can readily be prepared by methods known <u>per se</u>.

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#### **EXAMPLE 1**

5-(4-Chloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a mixture of benzyloxyacetaldehyde (8.3 g, 0.06 mol) in benzene (80 mL) was added 1-methoxy-3-trimethylsilyloxy-1,3-butadiene (10.6 g, 0.06 mol). The reaction mixture was stirred under nitrogen for 15 min., cooled to 0 °C and a solution of 0.5 M zinc chloride (55 ml, 0.03 mol) was added dropwise. The reaction mixture was allowed to warm to room temperature over 16 h and evaporated in vacuo. The resultant oil was diluted with ethyl acetate (100 ml), washed with 1N hydrochloric acid (3 x 50ml), saturated sodium bicarbonate (3 x 50 ml), brine (3 x 50 ml), dried (MgSO<sub>4</sub>) and evaporated in vacuo. The resulting oil was subjected to flash chromatography using a mixture of ethyl acetate/hexanes (1:2) as eluant. Pure fractions were collected affording after evaporation in vacuo 7.1 g

(60 %) of benzyloxy-methyl-2,3-dihydro-pyran-4-one as an oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.39 - 7.31 (m, 6H), 5.42 (dd, J = 6,1 Hz, 1H), 4.61 (d, J = 3 Hz, 1H), 4.57 (m, 1H), 3.70 (m, 2H), 2.74 (dd, J = 17)

Hz, 14 Hz, 1H), 2.41 (ddd, J = 17 Hz, 2 Hz, 1 Hz, 1H).

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The above 2,3-dihydro-pyran-4-one (7.1 g, 0.032 mol) and 10 % palladium on carbon (0.4 g) in ethyl acetate (50 ml) were placed in a Parr bomb shaker and hydrogenated at 30 psi. The reaction mixture was shaken for 2 h, at-which time TLC analysis (methanol/dichloromethane 1.9) indicated the reaction was complete. The reaction mixture was filtered through a pad of Celite and the volatiles evaporated in vacuo. The residue was

subjected to flash column chromatography using ethyl acetate as eluant. Pure fractions were collected affording after evaporation in vacuo 3.0 g (75 %) of 2-hydroxymethyl-tetrahydro-pyran-4-one as an oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  4.36 - 4.29 (m, 1H), 3.77 - 3.66 (m, 3H), 3.61 - 3.54 (m, 1H), 2.65 - 2.43 (m, 2H), 2.34 - 2.27 (m, 2H), 2.04 (bs, 1H, CH<sub>2</sub>O*H*).

The above tetrahydro-pyran-4-one (1.90 g, 0.015 mol), *tert*-butyl cyanoacetate (2.7 g, 0.019 mol), sulfur (0.51 g, 0.016 mol) and morpholine (2.55 ml, 0.03 mol) were dissolved in absolute ethanol (20 ml), and heated to 50 °C for 16 h. The reaction mixture was cooled, filtered and the filtrate evaporated <u>in vacuo</u>. The resultant oil was dissolved in ethyl acetate (50 ml), washed with water (2 x 50 ml), brine (2 x 50 m) and dried (MgSO<sub>4</sub>). The solvent was evaporated <u>in vacuo</u> and the residue was subjected to flash column chromatography using ethyl acetate/hexanes (1:1) as eluant. Pure fractions were collected affording after evaporation <u>in vacuo</u> 3.7 g (90 %) of 2-amino-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 4.64 (s, 2H), 3.80 - 3.67 (m, 3H), 2.77 - 2.72 (m, 1H), 2.57 - 2.53 (m, 1H), 1.54 (s, 9H).

The above carboxylic acid *tert*-butyl ester (1.0 g, 3.5 mmol), 4-chloro-1,3-dioxo-1,3-dihydro-isoindol (0.67 g, 3.7 mmol) and triphenylphosphine (1.01 g, 3.9 mmol) were dissolved in dry tetrahydrofuran (30 ml) and cooled to 0 °C under a nitrogen atmosphere. Diisopropyl azodicarboxylate (DIAD) (0.62 ml, 3.9 mmol) was added dropwise at 0 °C and the solution allowed to stir overnight, slowly warming to room temperature. The volatiles were evaporated <u>in vacuo</u> and the resultant solid dissolved in ethyl acetate (50 ml). The organic phase was washed with brine (3 x 50 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated <u>in vacuo</u>. The residue was subjected to flash column chromatography (300 ml silicagel) using a mixture of ethyl acetate/hexanes (1:3) as eluant. Semi pure fractions were collected affording after evaporation <u>in vacuo</u> 0.7 g which was trituated with diethyl ether. The solid was filtered off and

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washed with diethyl ether and dried <u>in vacuo</u> affording 0.13 g (27 %) of 2-amino-5-(4-chloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid. The filtrate was evaporated <u>in vacuo</u>. The residue (0.48 g) was subjected to flash column chromatography (300 ml silicagel) using a mixture of ethyl acetate/hexanes (1:3) as eluant. Pure fractions were collected affording after evaporation <u>in vacuo</u> an additional 0.36 g (23 %) of 2-amino-5-(4-chloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

To the above 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert butyl ester (0.36 g, 0.8 mmol) dissolved in tetrahydrofuran (20 ml) was added a mixture of imidazol-1-yl-oxo-acetic acid tert butyl ester (0.31 g, 1.6 mmol) in tetrahydrofuran (3.4 ml) under nitrogen. The reaction mixture was allowed to stir at room temperature for 18 hours. An additional portion of imidazol-1-yl-oxo-acetic acid tert butyl ester (0.3 g, 1.6 mmol) in tetrahydrofuran (2 ml) was added. The reaction mixture was allowed to stir at room temperature for an additional 60 h. The reaction mixture was poured into water (50 ml) and extracted with ethyl acetate (3 x 50 ml). The combined organic phases were washed with brine (3 x 50 ml) dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the organic phase evaporated in vacuo. The residue (0.5 g) was purified by column chromatography (300 ml silicagel) using a mixture of ethyl acetate/heptane (1:2) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 0.36 g (80 %) of 2-(tert-butoxyoxalyl-amino)-5-(4-chloro-1,3-dioxo-1,3-dihydroisoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

The above di-*tert*-butyl ester (0.3 g, 0.52 mmol) was dissolved in dichloromethane (1.2 ml) and trifluoroacetic acid (0.5 ml) was added. The reaction was stirred at room temperature for 18 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with a mixture of diethyl ether and heptane (1:1) (5 ml). The precipitate was filtered off, washed with heptane and diethyl ether, dried <u>in vacuo</u> at 50 °C for 18 h which afforded 200 mg (69 %) of the <u>title compound</u> as a solid.

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M.p.: > 250 °C

Calculated for C<sub>19</sub>H<sub>13</sub>N<sub>2</sub>ClO<sub>8</sub>S;

C, 49.09 %; H, 2.82 %; N, 6.03 %. Found:

5 C, 48.79 %; H, 2.79 %; N, 5.89 %.

#### **EXAMPLE 2**

10 <u>5-(4,5,6,7-Tetrachloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid</u>

In a 4 ml scintillating vial, a solution of tetrachloro phthalimide (148 mg, 0.52 mmol) in N,N-dimethylformamide (2.0 ml) was heated to 100°C for 10 minutes and treated with potassium hydride (55 mg, 0.48 mmol, 35 % w/w dispersion in mineral oil). The resulting mixture was stirred until gas generation ended, 2-(tert-butoxyoxalyl-amino)-5-(4-nitro-benzenesulfonyl-oxymethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (151 mg, 0.25 mmol) and 18-crown-6 ether (31 mg, 0.12 mmol) were added. The solution was flushed with nitrogen gas before being stirred at 80°C for 25 h. The volatiles were evaporated in vacuo and the residue purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (5:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 39 mg (23 %) of 2-(tert-butoxyoxalyl-amino)-5-(4,5,6,7-tetrachloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  12.50 (s, 1H), 4.80 (d, J = 16, 1H), 4.67 (d, J = 14, 1H), 4.14-3.99 (m, 2H), 3.84( d, J = 9, 1H), 2.99 (d, J = 17, 1H), 2.70 (dd, J = 17, 5, 1H), 1.60 (s, 9H), 1.56 (s, 9H).

HPLC (254.4 nm) R<sub>t</sub>=5.80 min, 95%.

In a 25 ml round bottom flask, 2-(*tert*-butoxyoxalyl-amino)-5-(4,5,6,7-tetrachloro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (39 mg, 0.06 mmol) was dissolved in 20 % trifluoroaceetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring for 24 h. A precipitate was filtered off and washed with diethyl ether, affording after drying 29 mg (90 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.32 (s, 1H), 4.76 (d, J = 16, 1H), 4.59 (d, J = 14, 1H), 4.0-3.6 (m partially obscured by water, 3H), 3.1 (d partially obscured by water, J = 17, 1H), 2.61 (dd partially obscured by DMSO, J = 20, 11, 1H).

HPLC (254.4 nm) R<sub>t</sub>=4.15 min, 75 %.

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#### **EXAMPLE 3**

5-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 4-hydroxyphthalic acid (0.25 g, 1.37 mmol) in anhydrous N,N-dimethylformamide (3 ml) under nitrogen was added sodium hydride (0.22 g, 5.48 mmol). The solution was stirred for 5 minutes and then methyl iodide (0.68 ml) was added and continued stirring for 3 hours. Several drops of water were added to quench the reaction and the mixture was concentrated in vacuo. The crude material was partitioned between ethyl acetate (40 ml) and water (10 ml). The layers were

separated and the organic layer washed with brine (2 x 10 ml), dried(Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The resulting oil was dissolved in methanol (8 ml) and 1N sodium hydroxide (4 ml) was added. The reaction was stirred at ambient temperature for 24 h., after which LC-MS indicated only partial hydrolysis. The material was reconstituted in methanol (5 ml) and treated with of sodium hydroxide (0.12 g, 3.0 mmol) dissolved in water (1 ml). The reaction mixture was stirred for 48 h., at which time a precipitate had formed. The mixture was acidified with 6N hydrochloric acid until pH = 1, causing the solution to become homogeneous. The reaction was concentrated <u>in vacuo</u> and the residue partitioned between ethyl acetate (30 ml) and 0.5N hydrochloric acid (10 ml). The layers were separated and the organic layer concentrated <u>in vacuo</u> to give 100 mg (51 %) of 4-methoxy-phthalic acid as a solid.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ 7.83 (d, J = 8, 1H), 7.10-7.06 (m, 2H), 3.87 (s, 3H).

LC-MS: R<sub>t</sub>=1.45 min,  $[M+H]^{+}$  = 197.1

A solution of 4-methoxy-phthalic acid (0.10 g, 0.51 mmol), 1-hydroxy-20 benzotriazole (0.15 g, 1.1 mmol), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.22 g, 1.1 mmol), and triethylamine (0.35 ml, 2.5 mmol) was prepared in distilled acetonitrile (4 ml) under nitrogen. 2-Amino-5-aminomethyl-4,7-dihydro-5H-thieno-[2,3-c]pyran-3-carboxylic acid tert-butyl ester (0.11 g, 0.39 mmol) was added in small portions and the reaction was stirred at ambient temperature for 18 h., and then 25 concentrated in vacuo. The crude mixture was diluted in ethyl acetate (30 ml) and washed with 1% hydrochloric acid (5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel chromatography using a 10 % mixture of 30 ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated in vacuo to give 54 mg (31 %) of 2-amino-5-

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(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno-[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  7.76 (d, J = 8, 1H), 7.32 (s, 1H), 7.14 (d, J = 8, 1H), 4.62-4.48 (m, 2H), 4.00-3.72 (m, 3H), 3.91 (s, 3H), 2.86 (d, J = 17, 1H), 2.55 (dd, J = 17, 10, 1H), 1.49 (s, 9H).

To a solution of the above 2-amino-5-(5-methoxy-1,3-dioxo-1,3dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno-[2,3-c]pyran-3carboxylic acid tert-butyl ester (54 mg, 0.12 mmol) in distilled dichloromethane (3 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid tert-butyl ester (0.25 g, 0.36 mmol) and triethylamine (50 ul, 0.36 mmol). The reaction was stirred for 4 h., concentrated in vacuo and the residue reconstituted in ethyl acetate (20 ml). The organic layer was washed with 1% hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The organic phase was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel chromatography using a 5% mixture of ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated in vacuo to give 56 mg (81%) of 2-(tertbutoxyoxalyl-amino)-5-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2ylmethyl)-4,7-dihydro-5H-thieno-[2,3-c]pyran-3-carboxylic acid tert-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.48 (s, 1H), 7.75 (d, J = 8, 1H), 7.32 (d, J = 2, 1H), 7.15 (dd, J = 8, 2, 1H), 4.78 (d, J = 15, 1H), 4.65 (d, J = 15, 1H), 4.03-3.75 (m, 3H), 3.91 (s, 3H), 2.95 (d, J = 17, 1H), 2.66 (dd, J = 17, 9, 1H), 1.58 (s, 9H), 1.54 (s, 9H). *APCI-MS*: [M+H]<sup>+</sup> = 574

The above 2-(*tert*-butoxyoxalyl-amino)-5-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno-[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (55 mg, 0.096 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (4 ml). The reaction

was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 17 mg (40%) of the <u>title compound</u> as a solid.

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<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.32 (s, 1H), 7.81 (d, J = 8 , 1H), 7.40 (d, J = 2, 1H), 7.31 (dd, J = 8 , 2, 1H), 4.75 (d, J = 15, 1H), 4.56 (d, J = 15, 1H), 3.92 (s, 3H), 3.91-3.69 (m, 3H), 2.98 (d, J = 17, 1H), 2.57 (dd, J = 17, 9, 1H).

10 APCI-MS: [M-H] = 459

HPLC (254.4nm): R<sub>t</sub>=3.36 min, 98%

#### **EXAMPLE 4**

OH OH

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5-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(*tert*20 butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester was prepared in a similar way as described in Example 1.

To a solution of the above benzylether (0.7 g, 1.08 mmol) in ethyl acetate (50 ml) was added 10 % palladium on carbon (0.2 g). The mixture was hydrogenated at 1 atm. for 5 h, filtered and the volatiles evaporated in vacuo. The residue (0.6 g) was purified by column chromatography (500 ml silicagel) using a mixture of ethyl acetate/heptane (1:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 0.4 g (67 %) of 2-(tert-butoxyoxalyl-amino)-5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-

30 carboxylic acid *tert*-butyl ester as an oil.

TLC:  $R_f = 0.2$  (ethyl acetate/heptane 1:1)

The above di-*tert*-butyl ester (0.4 g, 0.72 mmol) was dissolved in 25 % trifluoroacetic acid in dichloromethane (25 ml). The reaction was stirred at room temperature for 18 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with diethyl ether (5 ml). The precipitate was filtered off, washed with heptane and diethyl ether, dried <u>in vacuo</u> at 50 °C for 18 h which afforded 230 mg (72 %) of the <u>title compound</u> as a solid.

M.p.: > 250  $^{\circ}$ C;

Calculated for C<sub>19</sub>H<sub>14</sub>N<sub>2</sub>O<sub>9</sub>S, 0.5 x H<sub>2</sub>O;

10 C, 50.11 %; H, 3.32 %; N, 6.15 %. Found: C, 50.06 %; H, 3.17 %; N, 5.98 %.

#### **EXAMPLE 5**

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<u>5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid</u>

5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(*tert*butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.7 g, 1.08 mmol) (prepared in a similar way as described in Example 1) was dissolved in 25 % trifluoroacetic acid in dichloromethane (25 ml). The reaction was stirred at room temperature for 18 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with diethyl ether (25 ml). The precipitate was filtered off, washed with diethyl ether and dried <u>in vacuo</u> at 50 °C for 3 hours which afforded 400 mg (69 %) of the title compound as a solid.

M.p.: 194 - 196 °C;

30 Calculated for  $C_{26}H_{20}N_2O_9S$ , 1 x  $H_2O$ , 0.6 x  $CF_3COOH$ ;

C, 52.44 %, H, 3.66 %, N, 4.50 %. Found: C, 52.33 %, H, 3.65 %, N, 4.62 %.

#### **EXAMPLE 6**

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5-(4-Fluoro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

10 Prepared in a similar way as described in Example 1.

M.p.: > 250 °C; Calculated for  $C_{19}H_{13}FN_2O_8S$ , 1 x  $H_2O$ ; C, 48.93 %; H, 3.24 %; N, 6.01 %. Found: C, 48.90 %; H, 3.15 %; N, 5.86 %.

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# EXAMPLE 7

5-(1,3-Dioxo-1,3-dihydro-benzo[f]isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

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In-a-4-ml-scintillating-vial, a solution of benzo[f]isoindole-1,3-dione (145 mg, 0.74 mmol) in N,N-dimethylformamide (2.0 ml) was treated with potassium hydride (55 mg, 0.48 mmol, 35 % w/w dispersion in mineral oil).

The resulting mixture was stirred until gas generation ended and the resulting precipitate was filtered off and washed with dichloromethane which afforded 121 mg (69 %) of benzo[f]isoindole-1,3-dione potassium salt as a solid.

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 $^{1}\text{H NMR}$  (300 MHz, D<sub>2</sub>O)  $\delta$  8.00-7.87 (m, 4H), 7.62 (s, 2H).

In a 4 ml scintillating vial, the above potassium salt (121 mg, 0.5 mmol) in N,N-dimethylformamide (1.5 ml) was treated with 18-crown-6 ether (34 mg, 0.13 mmol) and 2-(*tert*-butoxyoxalyl-amino)-5-(4-nitrobenzene-sulfonyloxymethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (148 mg, 0.25 mmol). The solution was flushed with nitrogen gas before being stirred at 80 °C for 7 h. The volatiles were evaporated <u>in vacuo</u> and the residue purified by silica gel chromatography using a mixture of ethyl acetate/dichloromethane (1:49) as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 85 mg (57 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-benzo[*f*]isoindole-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.52 (s, 1H), 8.37 (s, 2H), 8.08 (m, 2H), 7.72 (m, 2H), 4.84-4.65 (m, 2H), 4.16-3.90 (m, 3H), 3.02 (d, J = 17, 1H), 2.73 (dd, J = 17, 10, 1H), 1.61 (s, 9H), 1.58 (s, 9H).

In a 25 ml round bottom flask the above 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-benzo[*f*]isoindole-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (85 mg, 0.14 mmol) was dissolved in 20 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring for 24 h. The precipitate was filtered off and washed with diethyl ether, affording after drying 62 mg (90 %) of the <u>title compound</u> as a solid.  $^1$ H-NMR (300-MHz, DMSO-d<sub>6</sub>)  $\delta$  12.32 (s, 1H), 9.02 (s, 2), 4.81-4.59 (m, 2H), 3.97-3.81 (m partially obscured by water, 3H), 3.08 (d, J = 18, 1H), 2.74-2.53 (m partially obscured by DMSO, 1H).

#### **EXAMPLE 8**

5-(5-Acetylamino-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

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To a solution of N-(1,3-dioxo-2,3-dihydro-1H-isoindol-5-yl)acetamide (51 mg, 0.25 mmol) in N,N-dimethylformamide (1.5 ml) under nitrogen at room temperature was added potassium hydride (35 wt.% dispension in mineral oil, 29 mg, 0.25 mmol). The solution was stirred at room temperature for 3 hours. A solid precipitated during this period. 2-(tert-Butoxyoxalyl-amino)-5-(4-nitro-benzene-sulfonyloxymethyl)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (100 mg. 0.17 mmol) was added to the suspension and the solution was stirred at ,80 °C for 12 h. The solvent was evaporated in vacuo, the resulting residue purified by silica gel chromatography using a gradient of ethyl acetate/hexane (10-25%) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 50 mg (50 %) of 5-(5-acetylamino-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(tert-butoxyoxalyl-amino)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  12.53 (s, 1H), 8.03 (d, 1H, J = 1.5 Hz), 7.91 (dd, 1H, J= 7.8, 1.8 Hz) 7.83 (d, 1H, J = 8.1 Hz), 7.45 (s, 1H), 4.80 (d, 1H, J = 16)Hz), 4.66 (d, 1H, J = 16 Hz), 4.03 (m, 2H), 3.83 (q, 1H, J = 15 Hz), 2.98 (d, 1H, J = 9 Hz), 2.64-2.78 (m, 1H), 2.27 (s, 3H), 1.62 (s, 9H), 1.57 (s, 9H).

To a mixture of trifluoroacetic acid/dichloromethane (2 ml, 1:1) at room temperature was added the above 5-(5-acetylamino-1,3-dioxo-1,3-

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dihydro-isoindol-2-ylmethyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (40 mg, 0.067 mmol). The solution was stirred for 5 h. at which time the solvent was removed <u>in vacuo</u>. The residue was washed with dichloromethane, filtered off, and dried <u>in vacuo</u> which afforded 23 mg (70 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 12.32 (s, 1H), 10.58 (s, 1H), 8.21 (s, 1H) 7.84 (s, 2H), 4.76 (d, 1H, J = 15 Hz), 4.58 (d, 1H, J = 15 Hz), 3.80-4.00 (m, 3H), 3.00 (d, 1H, J = 17 Hz), 2.58-2.73 (m, 1H), 2.13 (s, 3H).

10 MS: 488 (M+1).

### **EXAMPLE 9**

15 <u>5-(4-Acetylamino-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid</u>

The <u>title compound</u> was prepared in a similar way as described for Example 8.

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 12.32 (s, 1H), 9.76 (s, 1H), 8.45 (d, 1H, J = 8.4 Hz) 7.79 (t, 1H, J = 8.4 Hz), 7.58 (d, 1H, J = 8.4 Hz), 4.77 (d, 1H, J = 15 Hz), 4.58 (d, 1H, J = 15 Hz), 3.68-3.94 (m, 3H), 3.02 (d, 1H, J = 16 Hz), 2.55-2.78 (m, 1H), 2.20 (s, 3H). MS: 488 (M+1).

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#### EXAMPLE 10

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# N O O S O OH

# 5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

In a 4-ml scintillating vial, a solution of 2-amino-5-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (148 mg, 0.5 mmol) in tetrahydrofuran (1.0 ml) was treated with a solution of pyrazine phthtalic acid anhydride (85 mg, 0.56 mmol) in tetrahydrofuran (1.0 ml) and N,N-dimethylformamide (0.5 ml). The reaction mixture was allowed to stir at room temperature for 1 h. Diisopropylethylamine (220 µl, 0.13 mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (121 mg, 0.6 mmol) were then added. The reaction mixture was shaken vigorously for 10 seconds before being stirred at room temperature for 14 h. The volatiles were evaporated in vacuo and the residue purified by silica gel chromatography using a mixture of dichloromethane/ethyl acetate (3:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 25 mg (12 %) of the 2-amino-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) d 8.97 (s, 2H), 4.62-4.49 (m, 2H), 4.21-4.04 (m, 2H), 3.94 (dd, J = 14, 4, 1H), 2.91 (d, J = 17, 1H), 2.63 (dd, J = 17, 10, 1H), 1.68 (s, 9H).

In a 4 ml scintillating vial a solution of the above 2-amino-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (25 mg, 0.06 mmol) in tetrahydrofuran (3 ml) was treated with midazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.36 mmol). After stirring for 3 hours at room temperature the reaction solution was concentrated to dryness <u>in vacuo</u>. The residue was purified by silica gel chromatography using a mixture-of hexanes/ethyl-acetate (3:1) as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 31 mg (95 %) of 2-(*tert*-butoxyoxalyl-

amino)-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.49 (s, 1H), 8.96 (s, 2H), 4.80-4.61 (m, 2H), 4.21-4.04 (m, 2H), 3.96 (dd, J = 14, 4, 1H), 3.03 (d, J = 16, 1H), 2.70 (dd, J = 17, 10, 1H), 1.60 (s, 9H), 1.59 (s, 9H).

In a 25 ml round bottom flask the above 2-(*tert*-butoxyoxalyl-amino)-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyrazin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester, (31 mg, 0.06 mmol) was dissolved in 20 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring for 24 h. A precipitate was filtered off and washed with diethyl ether, affording after drying 22 mg (90 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 9.02 (s, 2), 4.81-4.59 (m,

5 2H), 3.97-3.81 (m partially obscured by water, 3H), 3.08 (d, J = 18, 1H), 2.74-2.53 (m partially obscured by DMSO, 1H).

HPLC (254.4 nm) R<sub>t</sub>=2.97 min, 89%. MS (APCI) [M-H] 432.4

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#### **EXAMPLE 11**

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7-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

A solution of furo[3,4-b]pyridine-5,7-dione (86.1 mg, 0.58 mmol) and of 2-(tert-butoxyoxalyl-amino)-7-aminomethyl-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (194 mg, 0.47 mmol) in acetonitrile (2.0 ml) was stirred for 10 min. at room temperature. 1-(3dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (89.2 mg, 0.564 mmol) and triethylamine (198 µl, 1.41 mmol) were added and the mixture was stirred at room temperature for 20 h. The volatiles were removed in vacuo and the crude product dissolved in dichloromethane (60 ml) and washed with water (3 x 30ml). The organic layer was dried (MgSO<sub>4</sub>). 10 filtered and the solvent removal in vacuo. The residue (338 mg) was purified by column chromatography on silica gel utilizing a mixture of hexane/ethyl acetate (90/10 to 50/50) as gradient which afforded after evaporation of the solvent in vacuo 85 mg (33 %) of 2-(tert-butoxyoxalylamino)-7-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-4,7-15 dihyd-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>),  $\delta$  9.00 (d, J = 4.8, 1H), 8.21 (d, J = 7.5, 1H), 7.64 (dd, J = 4.8, J = 6.8, 1H), 5.12 (d, J = 7.2, 1H), 4.24-4.1 (m, 2H), 3.97-3.91 (m, 1H), 3.75 (m, 1H), 2.90 (m, 1H), 1.29 (s, 9H), 1.27 (s, 9H). MS: 544 (M+1).

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The above 2-(*tert*-butoxyoxalyl-amino)-7-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-4,7-dihyd-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (47.4 mg, 0.087 mmol) was stirred in 50% trifluoroacetic acid in dichloromethane (2 ml) at room temperature for 5 h. The solvent was removed <u>in vacuo</u> and the residue was washed with diethyl ether (4 x 3.0 ml) and dried which afforded 26.5 mg (70 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (400MHz, CD<sub>3</sub>OD): δ 8.96 (d, J = 5, 1H), 8.30 (d, J = 7.6, 1H), 7.79 (dd, J = 5.2, J = 5.2, 1H), 5.10 (d, J = 6.4, 1H), 4.16 (m, 2H), 3.96 (dd, J = 3.2, J = 3.6, 1H), 3.78 (m, 1H), 2.95 (m, 2H). MS: 432 (M+1).

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5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

Pyrrolo[3,4-b]pyridine-5,7-dione (74.2 mg, 0.5 mmol) was stirred with sodium hydride (60% dispersion in mineral oil, 20.04 mg, 0.5 mmol) in N,N-dimethylformamide (4.0 ml) at room temperature under inert atmosphere. 2-(*tert*-Butoxyoxalyl-amino)-5-(4-nitro-benzene-sulfonyloxymethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (198 mg, 0.33 mmol) was added to the sodium salt formed and the reaction was stirred at 80 °C for 20 h. The solvent was removed <u>in vacuo</u> and the crude product was purified by preparative TLC (hexane:ethyl acetate 50:50) which afforded 58 mg (21 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  9.00 (d, J = 5, 1H), 8.20 (d, J = 7.5, 1H), 7.65 (dd, J = 5, J = 5, 1H), 4.80 (d, J = 14.7, 1H), 4.66 (d, J = 14.7, 1H), 4.10 (m, 2H), 3.91 (d, J = 13.2, 1H), 3.02 (d, J = 16.5, 1H), 2.70 (m, 1H), 1.61 (s, 9H), 1.58 (s, 9H).

MS: 544 (M+1).

The above 2-(*tert*-butoxyoxalyl-amino)-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-b]pyridin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (46.4 mg, 0.09 mmol) was stirred in 20 % trifluoroacetic acid in dichloromethane (3.0 ml) at room temperature for 2 h. The volatiles were removed <u>in vacuo</u> and the residue was washed with diethyl ether (5 x 3 ml) affording 37 mg (99 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H-NMR-(300-MHz, GDCl<sub>3</sub>): δ 8:96 (d, J = 5.4, 1H), 8.20 (d, J = 7.7, 1H), 30 7.64 (m, 1H), 4.77 (d, J = 14.7, 1H), 4.61 (d, J = 14.7, 1H), 4.07 (m, 2H), 3.86 (d, J = 10.5, 1H), 3.12 (d, J = 17.4, 1H), 2.77-2.68 (m, 2H).

MS: 432 (M+1).

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#### **EXAMPLE 13**

5-(5,7-Dioxo-5,7-dihydro-pyrrolo[3,4-c]pyridin-6-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of pyrrolo[3,4-c]pyridine-1,3-dione (74 mg, 0.50 mmol) in N,N-dimethylformamide (1 ml) under nitrogen at room temperature was added potassium hydride (35 wt.% dispersion in mineral oil, 57 mg, 0.50 mmol). The solution was stirred at room temperature for 3 hours. A solid precipitated during this period. 18-Crown-6 (33 mg, 0.13 mmol) and 2-(tert-butoxyoxalyl-amino)-5-(4-nitro-benzene-sulfonyloxymethyl)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (150 mg, 0.25 mmol) were then added. The solution was stirred at 80°C for 12 h and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/hexane (10-25%) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 93 mg (68 %) of 2-(tert-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester as a solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  12.49 (s, 1H), 9.20 (s, 1H), 9.11 (d, 2H, J = 4.8 Hz) 7.80 (d, 2H, J = 4.8 Hz), 4.80 (d, 1H, J = 16 Hz), 4.66 (d, 1H, J = 16 Hz), 4.00-4.18 (m, 2H), 3.70-3.95 (m, 1H), 3.01 (d, 1H, J = 17 Hz), 2.64-2.78(m, 1H), 1.60 (s, 9H), 1.59 (s, 9H).

To a mixture of trifluoroacetic acid/dichloromethane (1 ml, 1:1) at room temperature was added the above 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (29 mg, 0.053 mmol). The solution was stirred for 5 h. and the solvent evaporated <u>in vacuo</u>. The

residue was washed with dichloromethane afford after drying <u>in vacuo</u> 22 mg (96 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 12.32 (s, 1H), 9.15 (s, 1H), 9.11 (d, 2H, J = 4.8 Hz) 7.92 (d, 2H, J = 4.8 Hz), 4.76 (d, 1H, J = 15 Hz), 4.58 (d, 1H, J = 16 Hz), 3.75-4.00 (m, 4H), 3.04 (d, 1H, J = 17 Hz). MS: 432 (M+1).

# **EXAMPLE 14**

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5-(5-Nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

In a 4-ml scintillating vial, a solution of 2-amino-5-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (58 mg, 0.2 mmol) in tetrahydrofuran (2.0 ml) was treated with 4-nitrophthalic acid (63 mg, 0.3 mmol), diisopropylethylamine (190 μl, 1.1 mmol), and 1,3-diisopropylcarbodiimide (120 μl, 0.77 mmol). The reaction mixture was shaken vigorously for 10 seconds before being stirred at 50 °C for 43 hours and at room temperature for 20 h. The solution was diluted with ethyl acetate (25 ml), washed with 0.5N aqueous hydrochloric acid (25 ml), saturated aqueous sodium bicarbonate (25 ml), and brine (25 ml). The organic layer was dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. Crude 2-amino-5-(5-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester was obtained as a solid and used immediately in the next step.



mmol). After stirring for 2 h. at room temperature the reaction mixture was concentrated to dryness in vacuo. The residue was purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (3:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo which afforded 30 mg (26 %) of 2-(tert-butoxyoxalyl-amino)-5-(5-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  12.47 (s, 1H), 8.71 (s, 1H), 8.64 (d, J = 8, 1H), 8.08 (d, J = 9, 1H), 4.79 (d, J = 14, 1H), 4.65 (d, J = 14, 1H), 4.21-3.97 (m, 2H), 3.89 (d, J = 12, 1H), 3.01 (d, J = 16, 1H), 2.83-2.61 (m, 1H), 1.63 (ds, 18H).

In a 25 ml round bottom flask, the above 2-(*tert*-butoxyoxalyl-amino)-5-(5-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (30 mg, 0.05 mmol) was dissolved in a mixture of 20 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring of 24 h. A precipitate was filtered off and washed with diethyl ether, affording after drying 22 mg (90 %) of the <u>title compound</u> as a solid.

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<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 8.63 (d, J = 8, 1H), 8.15 (d, J = 8, 1H), 4.76 (d, J = 16, 1H), 4.57(d, J = 16, 1H), 4.42-3.74 (m partially obscured by water, 3H), 3.04 (d partially obscured by water, J = 16, 1H), 2.61 (m partially obscured by DMSO, 1H).

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HPLC (254.4 nm) R<sub>t</sub>=3.40 min, 86%. MS (APCI<sup>+</sup>) [M+H] 407.6

5-(5-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

- To a solution of 4-hydroxyphthalic acid (0.45 g, 2.47 mmol) in anhydrous N,N-dimethylformamide (5 ml) under nitrogen was added chloromethyl methyl ether (1.13 ml, 14.8 mmol) and diisopropylethylamine (2.6 ml, 14.8 mmol). The reaction was stirred at ambient temperature for 18 h. and then concentrated in vacuo. The crude material was partitioned between ethyl acetate (50 ml) and water (15 ml). The layers were separated, the organic layer washed with water (3 x 10 ml), brine (2 x 10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>)
  - layer washed with water (3 x 10 ml), brine (2 x 10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The resulting oil was dissolved in ethanol (5 ml) and sodium hydroxide (0.12 g, 7.4 mmol) dissolved in water (1 ml) was added to the reaction. The solution was
- stirred at ambient temperature for 48 h. and then concentrated in vacuo affording 4-methoxymethoxy-phthalic acid di-sodium salt which was used without purification.
  - <sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD)  $\delta$  7.59 (d, J = 8, 1H), 7.06 (d, J = 3, 1H), 6.89 (dd, J = 8, 3, 1H), 5.18 (s, 2H), 3.42 (s, 3H).
- A solution of 4-methoxymethoxy-phthalic acid di-sodium salt (0.19 g, 0.70 mmol), 1-hydroxybenzotriazole (0.2 g, 3.6 equiv.), 1-(3-dimethylamino-propyl)-3-ethylcarbodiimide hydrochloride (0.28 g, 3.6 equiv.), and triethylamine (0.33 ml, 6 equiv.) was prepared in distilled acetonitrile (5 ml) under nitrogen. The mixture was stirred for 5 minutes before 2-amino-5-
- aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (115 mg, 0.40 mmol) was added in small portions. The reaction was stirred at ambient temperature for 18 h., then concentrated in vacuo. The crude mixture was diluted with ethyl acetate (30 ml) and washed with 1% hydrochloric acid (5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The argania layer was dried (Ne SO.) filtered, and the activate
- ml). The organic layer was dried(Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel

chromatography using a gradient of ethyl acetate/dichloromethane (5 to 10% gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo to give 44 mg (23 %) of 2-amino-5-(5-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  7.75 (d, J = 8, 1H), 7.48 (d, J = 2, 1H), 7.27 (dd, J = 8, 2, 1H), 5.26 (s, 2H), 4.60-4.46 (m, 2H), 3.99-3.71 (m, 3H), 3.47 (s, 3H), 2.85 (d, J = 17, 1H), 2.55 (dd, J = 17, 9, 1H), 1.48 (s, 9H).

- 10 To a solution of the above 2-amino-5-(5-methoxy-methoxy-1,3-dioxo-1,3dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester (44 mg, 0.095 mmol) in distilled dichloromethane (3 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid tert-butyl ester (56 mg, 0.29 mmol) and triethylamine (26 µl, 0.19 mmol). The reaction was stirred for 4 h., concentrated in vacuo and 15 reconstituted in ethyl acetate (20 ml). The organic layer was washed with 1% hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The resulting solution was dried(Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel chromatography using a 5 % mixture of ethyl acetate/dichloromethane as 20 eluant. Pure fractions were collected and the solvent evaporated in vacuo to give 35 mg (63 %) of 2-(tert-butoxyoxalyl-amino)-5-(5-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3c|pyran-3-carboxylic acid tert-butyl ester.
  - <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.50 (s, 1H), 7.75 (d, J = 8, 1H), 7.49 (d, J = 2, 1H), 7.28 (dd, J = 8, 2, 1H), 5.26 (s, 2H), 4.77 (d, J = 15, 1H), 4.64 (d, J = 15, 1H), 4.03-3.74 (m; 3H), 3.47 (s, 3H), 2.95 (d, J = 17, 1H), 2.65 (dd, J = 17, 9, 1H), 1.58 (s, 9H), 1.54 (s, 9H). *APCI-MS*: [M+H]<sup>+</sup> = 603.7

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The-above-2-(*tert*-butoxyoxalyl-amino)-5-(5-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (35 mg, 0.058 mmol) was dissolved in a

mixture of 50 % trifluoroacetic acid/dichloromethane (2.5 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and the residue evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 20 mg (77 %) of the <u>title compound</u> as a solid.

1H NMR (300 MHz, DMSO-de)  $\delta$  12.31 (s. 1H), 10.97 (s. 1H), 7.72 (d. J =

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 10.97 (s, 1H), 7.72 (d, J = 8, 1H), 7.18 (s, 1H), 7.10 (d, J = 8, 1H), 4.74 (d, J = 15, 1H), 4.58 (d, J = 15, 1H), 3.96-3.62 (m, 3H), 2.99 (d, J = 17, 1H), 2.60-2.50 (m, 1H, partially obscured by DMSO).

10 APCI-MS: [M-H] = 445.4 HPLC (254.4nm): R<sub>t</sub>=2.92 min, 95%

### EXAMPLE 16

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5-(4-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

- To a solution of 4-hydroxy-isobenzofuran-1,3-dione (195 mg, 1.2 mmol) in anhydrous N,N-dimethylformamide (4 ml) under nitrogen was added sodium hydride (61 mg, 1.56 mmol). The solution was stirred for 15 minutes and then methyl iodide (0.37 ml, 6.0 mmol) was added. The reaction was stirred for 48 h. and then quenched with saturated
- ammonium chloride. The mixture was concentrated in vacuo, diluted in ethyl acetate (20 ml) and the organic phase washed with 1N hydrochloric acid (5 ml) and brine (3 x 5 ml). The organic layer was dried(MgSO₄) and concentrated in vacuo. To the crude solid was added methanol causing a precipitate to form. The flask was cooled in an ice bath for 2 h. and the

solid filtered off, washed with methanol and dried <u>in vacuo</u> which afforded 0.1 g (47 %) of 4-methoxy-isobenzofuran-1,3-dione as a solid. <sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>)  $\delta$  7.95 (t, J = 8, 1H), 7.61 (d, J = 8, 1H), 7.58 (d, J = 8, 1H), 3.99 (s, 3H).

5 APCI-MS:  $[M+H]^+ = 179.1$ 

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A solution of 2-amino-5-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (122 mg, 0.43 mmol, prepared as described in Example 17) and 4-methoxy-isobenzofuran-1,3-dione (92 mg, 0.52 mmol) was prepared in distilled tetrahydrofuran (4 ml) under nitrogen. 1-hydroxybenzotriazole (87 mg, 0.65 mmol), 1-(3-dimethyl-aminopropyl)-3-ethylcarbodiimide hydrochloride (123 mg, 0.65 mmol), and triethylamine (0.29 ml, 2.15 mmol) were added. The reaction was stirred at ambient temperature for 18 h., then concentrated in vacuo. The crude mixture was diluted with ethyl acetate (25 ml) and washed with 1N hydrochloric acid (5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo to give 0.18 g (94 %) of 2-amino-5-(4-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 7.66 (t, J = 7, 1H), 7.43 (d, J = 7, 1H), 7.19 (d, J = 7, 1H), 4.59-4.46 (m, 2H), 4.06-3.72 (m, 3H), 4.00 (s, 3H), 2.87-2.81 (m, 1H), 2.60-2.51 (m, 1H), 1.48 (s, 9H).

To a solution of the above 2-amino-5-(4-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.18 g, 0.42 mmol) in distilled dichloromethane (5 ml) under nitrogen was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.25 g, 1.26 mmol) and triethylamine (0.23 ml, 1.68 mmol). The reaction was stirred for 12 h., concentrated in vacuo and reconstituted in ethyl acetate (25 ml). The organic layer was washed with 1N hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The resulting solution was dried(Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated

in vacuo. The crude material was purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (0 to 10 % gradient). Pure fractions were collected and the solvent evaporated in vacuo to give 195 mg (81 %) of 2-(tert-butoxyoxalyl-amino)-5-(tert-butoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxalyl-aminoxyoxaly

The above 2-(*tert*-butoxyoxalyl-amino)-5-(4-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-

- carboxylic acid *tert*-butyl ester (0.15 g, 0.26 mmol) was dissolved in a mixture of 50 % trifluoroacetic acid/dichloromethane (5 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and the residue evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 100 mg (83 %) of the <u>title compound</u> as a solid.
  - <sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 7.79 (t, J = 8, 1H), 7.48 (d, J = 8, 1H), 7.42 (d, J = 8, 1H), 4.74 (d, J = 15, 1H), 4.56 (d, J = 15, 1H), 3.95 (s, 3H), 3.91-3.79 (m, 2H), 3.69-3.63 (m, 1H), 2.98 (d, J = 17, 1H), 2.57 (dd, J = 17, 10, 1H).
- 25 *LC-MS*:  $R_t$ =1.26 min,  $[M+H]^{+}$  = 461.0 HPLC (254.4nm):  $R_t$ =3.10 min, 100 %

LC-MS:  $R_t$ =4.17 min,  $[M+H]^{+}$  = 573.2

### **EXAMPLE 17**

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# 5-(4-Nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

In a 50-ml round-bottom flask, a suspension of 2-amino-5-(1,3-dioxo-1,3dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester (2.00 g, 4.8 mmol) in absolute ethanol (20 ml) was flushed with nitrogen and sealed with a rubber septum. Hydrazine (0.5 ml, 15.9 mmol) was added, followed by an additional portion of absolute ethanol (20 ml) at room temperature. The reaction mixture was heated to 80 °C for 3.5 h., then allowed to stir at room temperature for 14 10. h. The precipitate was filtered off and washed with absolute ethanol. The filtrate was concentrated in vacuo leaving an oil, which was dissolved in dichloromethane (30 ml) and refiltered. The solvent was evaporated in vacuo affording 1.2 g (86 %) of 2-amino-5-aminomethyl-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid. 15 <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) d 5.92 (s, 2H), 4.64 (s, 2H), 3.68-3.60 (m, 1H), 2.98-2.74 (m, 3H), 2.56-2.44 (m, 1H), 1.54 (s, 9H). MS (APCI<sup>+</sup>) [M+H] 285.3

In a 4-ml scintillating vial, a solution of the above 2-amino-5-aminomethyl-20 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (63 mg, 0.2 mmol) in tetrahydrofuran (2.0 ml) was treated with 3-nitro-phthalic acid (66 mg, 0.3 mmol), diisopropylethylamine (190 µl, 1.1 mmol), and 1,3-diisopropyl-carbodiimide (120 µl, 0.77 mmol). The reaction mixture was shaken vigorously for 10 seconds before being stirred at 50°C for 43 25 hours and at room temperature for 20 h. The reaction mixture was diluted with ethyl acetate (25 ml) and washed with 0.5N aqueous hydrochloric acid (25 ml), saturated sodium bicarbonate (25 ml), and brine (25 ml). The organic layer was dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo affording crude 2-amino-5-(4-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-30 ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

In a 4 ml scintillating vial a solution of the above crude 2-amino-5-(4-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester in dichloromethane (3 ml) was treated with midazol-1-yl-oxo-acetic acid *tert*-butyl ester (147 mg, 0.75 mmol). After stirring for 2 h. at room temperature the reaction solution was concentrated to dryness <u>in vacuo</u>. The residue was purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (3:1) as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 30 mg (26%) of 2-(*tert*-butoxyoxalyl-amino)-5-(4-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

1 H NMR (300 MHz, CDCl<sub>3</sub>) 8 8.17 (d, *J* = 5, 1H), 8.11 (d, *J* = 6, 1H), 7.94

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  8.17 (d, J = 5, 1H), 8.11 (d, J = 6, 1H), 7.94 (t, J = 8, 1H), 4.80 (d, J = 14, 1H), 4.67 (d, J = 15, 1H), 4.16-3.97 (m, 3H), 3.88 (d, J = 10, 1H), 3.01 (d, J = 16, 1H), 2.70 (dd, J = 16, 10, 1H), 1.62 (s, 9H), 1.59 (s, 9H).

In a 25 ml round bottom flask, the above 2-(*tert*-butoxyoxalyl-amino)-5-(4-nitro-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (30 mg, 0.05 mmol) was

20 dissolved in a mixture of 20 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring. After standing for 24 h. a precipitate was filtered off and washed with diethyl ether, affording after drying 22 mg (90 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.33 (s, 1H), 8.32 (d, *J* = 9, 1H), 8.20 (d, *J* = 9, 1H), 8.07 (t, *J* = 9, 1H), 4.77 (d, *J* = 14, 1H), 4.59 (d, *J* = 16, 1H), 4.00-3.65 (m partially obscured by water, 3H), 3.04 (d partially obscured by water, *J* = 16, 1H), 2.63 (dd partially obscured by DMSO, *J* = 17, 13,

HPLC (254.4 nm)  $R_1$ = 3.33 min, 100%.

30 MS (APCI<sup>+</sup>) [M+H] 391.6

1H).

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5-(4-(4-Chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

Under a nitrogen atmosphere, 4-(4-chloro-phenylsulfanyl)-6-methyl-pyrrolo[3,4-c]-1,3-dione (914 mg, 3.0 mmol), tributylphosphine (1.66 ml, 4.5 mmol) and 2-amino-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (855 mg, 3.0 mmol) were successively dissolved in dry benzene (90 ml). Solid azodicarboxylic dipiperidine (1.13 g, 4.5 mmol) was added under stirring at 0 °C to the solution. After stirring for 10 min, the reaction mixture was brought to room temperature and the stirring continued for 4 h. The mixture was cooled on ice, and additional portions of tributylphosphine (1.66 ml, 4.5 mmol) and azodicarboxylic dipiperidine (1.13 g, 4.5 mmol) were added. After stirring for 10 min, the reaction mixture was brought to room temperature and the stirring continued for 18 h. Heptane (30 ml) was added to the reaction and the precipitate filtered off (discard). After evaporation of the solvent the product was purified by flash chromatography to give 1.3 g (76 %) of 2-amino-5-(4-(4-chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-

Mp: 118 - 119° C;

carboxylic acid tert-butyl ester as an oil.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 1.55 (s, 9H), 2.50 (s, 3H), 2.50-2.65 (m, 1H), 2.85- 2.95 (m, 1H), 3.75-3.85 (m, 1H), 3.95- 4.05, (m, 2H), 4.50- 4.15 (m, 2H), 5.95 (bs, 2H), 7.30 (s, 1H), 7.40 (d, 2H), 7.55 (d, 2H).

pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-

To an ice cooled solution of 2-amino-5-(4-(4-chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (572 mg, 1 mmol) and dry triethylamine (2 ml) in dry tetrahydrofuran (10 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (588 mg, 3 mmol). After 10 min, the reaction mixture was brought to room temperature and the stirring continued for 18 h. The mixture was concentrated in vacuo and submitted to flash chromatography using a mixture of toluene/ethyl acetate (30:1) as eluant. Pure fraction were collected and the solvent evaporated in vacuo to give 360 mg (51 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(4-(4-chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

- 15 M.p.: 134 136° C; <sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 1.60 (s, 9H), 1.63 (s, 9H), 2.50 (s, 3H), 2.65-2.75 (m, 1H), 2.95- 3.05 (m, 1H), 3.75-3.90 (m, 1H), 4.00- 4.10, (m, 2H), 4.60- 4.85 (m, 2H), 7.30 (s, 1H), 7.40 (d, 2H), 7.55 (d, 2H), 12.50 (s, 1H).
- To 2-(*tert*-butoxyoxalyl-amino)-5-(4-(4-chloro-phenylsulfanyl)-6-methyl-1,3-dioxo-1,3-dihydro-pyrrolo[3,4-c]pyridin-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (324 g, 0.46 mmol) was added a mixture of trifluoroacetic acid (2.5 ml) and dichloromethane (7.5 ml). The mixture was stirred for 5 h, and added petroleum ether/ethyl acetate. The precipitate was isolated off and re-suspended in ethyl acetate. The <u>title compound</u> 136 mg (50 %) was isolated by filtration.

Mp: 239 - 240° C; Calculated for  $C_{25}H_{18}CIN_3O_8S_2$ , 0.75 x  $H_2O$ ; C, 49.92 %; H, 3.27 %; N, 6.99 %. Found:

30 ... C, 49.83 %; H, 3.16 %; N, 6.85 %. ....

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 2.48 (s, 3H), 2.65-2.75 (m, 1H), 2.95- 3.05 (m, 1H), 3.50-4.00 (m, 3H), 4.50- 4.90 (m, 2H), 7.50-7.68 (m, 5H), 12.30 (s, 1H).

### **EXAMPLE 19**

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5-(3-Imidazol-1-yl-2,5-dioxo-pyrrolidin-1-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

10 To a solution of 2-amino-5-aminomethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester (0.53 g, 1.86 mmol, prepared as described in Example 17) in tetrahydrofuran (10 ml) was added, maleic acid (0.24 g, 2.05 mmol) and diisopropylcarbodiimide (0.58 ml, 3.72 mmol). The reaction mixture was heated to reflux for 3 hours and 15 then allowed to cool to room temperature over an 18 hour period. The solvent was stripped off in vacuo and the residue diluted into ethyl acetate (50 ml). The organic phase was washed with saturated sodium bicarbonate (2 x 50 ml), 1 % hydrochloric acid (2 x 20 ml), brine (3 x 50 ml), dried(MgSO<sub>4</sub>), filtered, and the solvent evaporated in 20 vacuo affording an oil which was subjected to flash chromatography using a mixture of ethyl acetate/hexanes (6:4) as eluant. Pure fractions (R<sub>f</sub>=0.25) were collected and the solvent evaporated in vacuo to give 0.60 g (90 %) of 2-amino-5-(2,5-dioxo-2,5-dihydropyrrol-1-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid 25 tert-butyl ester as an oil.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  7.31 (d, J = 5.7, 1H), 6.63 (d, J = 5.4, 1H), 5.94 (bs, 2H), 4.67 (s, 2H), 3.93 (m, 1H), 3.82 (m, 2H), 2.89-2.83 (m, 1H), 2.69-2.60-(m, 1H), 1.54 (s, 9H).

MS: APCI (+): 365.2 (M+H);

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To a solution of the above 2-amino-5-(2,5-dioxo-2,5-dihydro-pyrrol-1-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (60 mg, 1.64 mmol) in tetrahydrofuran (2 ml) was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (50 mg, 2.46 mmol). The solution was stirred at room temperature for 48 h. The solvent was stripped off <u>in vacuo</u> and the resultant oil diluted in ethyl acetate (20 ml), washed with brine (3 x 25 ml), dried(MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was subjected preparative thin layer chromatography using a mixture of

methanol/dichloromethane (1:9) as eluant which afforded 25 mg (28 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(3-imidazol-1-yl-2,5-dioxo-pyrrolidin-1-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a mixture of diastereoisomers.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 7.65 (s, 1H), 6.94 (s, 1H), 5.92 (m, 1H), 5.22 (m, 1H), 4.68-4.53 (m, 2H), 4.00 (m, 3H), 3.71 (m, 1H), 3.47-3.38 (m, 1H), 3.03-2.87 (m, 1H), 2.61 (m, 1H), 1.60 (s, 9H), 1.54 (s, 9H). MS: APCI (+): 561.2 (M+H).

To the above 2-(*tert*-butoxyoxalyl-amino)-5-(3-imidazol-1-yl-2,5-dioxopyrrolidin-1-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl éster (25 mg, 0.05 mmol) was added a mixture of 20% trifluoroacetic acid in dichloromethane (2 ml). The reaction mixture was allowed to stir at room temperature for 2 h., at which time the mixture was concentrated <u>in vacuo</u>. The resultant solid was triturated with diethyl ether (2x) which afforded 13 mg (65 %) of the <u>title</u> compound as a solid.

 $^{1}$ H NMR (300 MHz, CD<sub>3</sub>OD) δ 9.15 (s, 1H), 7.78 (s, 1H), 7.63 (m, 1H), 5.75 (m, 1H), 4.69 (m, 2H), 4.46 (m, 1H), 3.85 (m, 2H), 3.66 (m, 1H), 3.02 (m, 1H), 2.83 (m, 1H), 2.64 (m, 1H), 2.46 (m, 1H).

30 MS: ESI (-): 447.4 (M-H).

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Oxalic acid 3-carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester methyl

To a solution of 2-amino-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-5 c]pyran-3-carboxylic acid tert-butyl ester (8.0 g, 28 mmol) in dry tetrahydrofuran (50 ml) was added midazol-1-yl-oxo-acetic acid tert-butyl ester (27.51 g, 0.14 mol) and triethylamine (3.93 ml, 0.14 mol). The reaction mixture was stirred at room temperature for 20 h. The volatiles were removed in vacuo and the crude product was dissolved in ethyl 10 acetate (300 ml) and washed with a saturated solution of sodium bicarbonate (3 x 100 ml), dilute hydrochloric acid (3 x 100 ml), water (3 x 100 ml) and brine (100 ml). The organic layer was dried(MgSO<sub>4</sub>), filtered and the solvent removed in vacuo affording a foam (16 g) which was purified on column chromatography on silica gel using a gradient of 15 hexane/ethyl acetate (90:10 to 50:50 gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo which afforded 11 g (91 %) of oxalic acid 2-amino-3-tert-butoxycarbonyl-4,7-dihydro-5Hthieno[2,3-c]pyran-5-ylmethyl ester tert-butyl ester as a solid. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  5.94 (s, 2H), 4.86 (d, J = 14.7, 1H), 4.77 (d, J = 14.4, 1H), 4.64 (m, 1H), 3.82-3.71 (m, 2H), 2.85 (d, J = 16.8, 1H), 2.68 20 (d, J = 10.5, 1H), 1.62 (s, 9H), 1.61 (s, 9H).MS: 414 (M+1).

A solution of the above oxalic acid 2-amino-3-*tert*-butoxycarbonyl-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester *tert*-butyl ester (8.3 g, 20.1 mmol) and potassium carbonate (1.7 g, 12.3 mmol) was stirred in methanol (80 ml) in presence of water (3 ml) at room temperature for 10 min., at which time TLC indicated reaction complete. Methanol was removed-in-vacuo and the crude product-was-dissolved-in-dichloromethane (300 ml) and washed with water (3 x 150 ml). The organic phase was dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in

vacuo. The residue was purified on flash chromatography on silica gel using a gradient of hexane/ethyl acetate (90:10 to 50:50 gradient) as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 0.65 g (9 %) of oxalic acid 2-amino-3-*tert*-butoxycarbonyl-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester methyl ester as a solid. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  4.86 (d, J = 15, 1H), 4.78 (d, J = 15, 1H), 4.00 (s, 3H), 3.82-3.70 (m, 3H), 2.86 (d, J = 17, 1H), 2.66 (dd, J = 10.2, J = 10.5, 1H), 1.62 (s, 9H). MS: 316 (M-55):

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To a solution of the above oxalic acid 2-amino-3-*tert*-butoxycarbonyl-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester methyl ester (160 mg, 0.43 mmol) in dry tetrahydrofuran (3.0 ml) was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (420.4 mg, 2.15 mmol) and triethylamine (120  $\mu$ l, 0.86 mmol). The resulting mixture was stirred at room temperature for 20 h. The solvent was evaporated in vacuo and the crude product was purified by flash chromatography on silica gel using a gradient of hexane/ethyl acetate (95.5 to 80:20 gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 173 mg (81 %) of oxalic acid 2-amino-3-*tert*-butoxycarbonyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester methyl ester as a solid. <sup>1</sup>H NMR (300MHz, CDCl<sub>3</sub>):  $\delta$  4.81 (dd, J = 14.7, J = 14.2, 2H), 4.40 (m, 2H), 4.00 (s, 3H), 2.96 (d, J = 15.3, 1H), 2.69 (dd, J = 10.8, J = 10.8, 1H), 1.61 (s, 9H), 1.57 (s, 9H).

25 MS: 388.3 (M-11).

The above oxalic acid 2-amino-3-*tert*-butoxycarbonyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl ester methyl ester (93.8 mg, 0.19 mmol) was stirred in 20 % trifluoroacetic acid in dichloromethane (2 ml) for 20 h. at room temperature. The solvent was removal-<u>in-vacuo</u>-which-afforded-73-mg-(95-%)-of-the-<u>title-compound</u>-as-a solid.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD):  $\delta$  4.76 (d, J = 5.7, 2H), 4.18 (d, J = 4.8, 2H), 3.97 (s, 3H), 2.99 (d, J = 16.2, 1H), 2.65 (d, J = 10.8, 1H). MS: 386 (M-1).

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#### **EXAMPLE 21**

Oxalic acid (3-carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl) ester

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To a solution of a mixture of 2-amino-5-hydroxymethyl-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester and 2-amino-7hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tertbutyl ester (1:4 estimated based on <sup>1</sup>H NMR) (200 mg, 0.70 mmol) and diisopropylethylamine (0.25 ml, 1.4 mmol) in dichloromethane (6.0 ml) cooled to 0 °C under nitrogen was added triethylchlorosilane (0.18 ml, 1.1 mmol). The solution was stirred at 0 °C for 5 min. and then stirred at room temperature for 15 min. The solution was washed with saturated sodium bicarbonate and brine, dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a 5 % mixture of ethyl acetate/hexane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 42 mg (16 %) of 2-amino-5triethylsilanyloxymethy-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (1) and 193 mg (69 %) of 2-amino-7-triethylsilanyloxymethy-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester **(2)**.

(2) <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  4.62 (s. 2H); 3.85-3.64 (m. 3H), 2.82 (dm. 1H, J =15 Hz), 2.49 (dd, 1H, J = 15, 11 Hz), 1.58 (s, 9H), 0.98 (t, 9 H, J = 7.8 Hz), 0.64 (q, 6H, J = 7.8 Hz).

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To a solution of 2-amino-7-triethylsilanyloxymethy-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (6.0 g, 15 mmol) in dichloromethane (10 ml) cooled to 0 °C under the nitrogen was added a solution of imidazol-1-yl-oxo-acetic acid tert-butyl ester (4.5 g, 18 mmol) in dichloromethane. The solution was stirred at 0 °C for 10 min. The reaction was quenched with water (1.0 ml). The solution was washed with brine and dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a 10% mixture of ethyl acetate/hexane as eluant. Pure fractions of two compounds were collected and the solvent evaporated in vacuo affording 4.5 g (56 %) of 2-(tertbutoxyoxalyl-amino)-7-triethylsilanyloxymethyl-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (A) as a solid and 50 mg of oxalic acid 3-(tert-butoxycarbonyl-2-(tert-butoxyoxalylamino)-4,7-dihydro-5H- thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (B) as a solid.

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  - (A) <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  12.53 (s, 1H), 4.85 (d, 1H, J = 12 Hz), 4.65 (d, 1H, J = 12 Hz), 3.90-3.60 (m, 3H), 2.94 (d, 1H, J = 15 Hz), 2.63 (dd, 1H, J = 15, 11 Hz), 1.63 (s, 9H), 1.61 (s, 9H), 0.98 (t, 9 H, J = 7.8 Hz), 0.64 (q, 6H, J = 7.8 Hz).
  - (B) <sup>1</sup>H NMR (CDCI<sub>3</sub>):  $\delta$  12.47 (s, 1H), 4.82 (q, 2H, J = 14 Hz), 4.43 (m, 2H), 4.01 (m, 1H), 2.97 (d, 1H, J = 14 Hz), 2.69 (dd, 1H, J = 19, 9 Hz), 1.63 (s, 9H), 1.61 (s, 9 H), 1.58 (s, 9H).

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To a solution of the above 2-(tert-butoxyoxalyl-amino)-7-triethylsilanyloxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tertbutyl ester (4.5 g, 8.5 mmol) in tetrahydrofuran (10 ml) at room

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temperature was added 0.5 N hydrochloric acid (2.0 ml). The solution was stirred at room temperature for 0.5 h. Ethyl acetate (100 ml) was added and the resulting solution was washed with saturated sodium bicarbonate, brine, dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a 10 % mixture of ethyl acetate/hexane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 3.0 g (84 %) of 2-(tert-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H NMR (CDCl<sub>3</sub>): δ 12.53 (s, 1H), 4.86 (d, 1H, J = 12 Hz), 4.60 (d, 1H, J = 12 Hz), 3.85-3.65 (m, 3H), 2.85 (d, 1H, J = 15 Hz), 2.65 (dd, 1H, J = 15, 11 Hz), 1.63 (s, 9H), 1.61 (s, 9H).

To a solution of the above 2-(tert-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (3.0 g, 7.1 mmol) in dichloromethane (10 ml) at room temperature was added pyridine (2.5 ml, 28.5 mmol) and 4-nitrobenzenesulfonyl chloride (4.7 g, 21.4 mmol). The solution was heated to 50 °C and stirred for 4.5 h. The solution was cooled to room temperature and washed with 0.5 N hydrochloric acid, saturated sodium bicarbonate, brine, dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/hexane (0-100 %) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 3.6 g (84 %) of 2-(tert-butoxyoxalyl-amino)-7-(4-nitrobenzenesulfonyloxymethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester as a solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  12.40 (s, 1H), 8.43 (d, 2H, J = 9.0 Hz), 8.17 (d, 2H, J = 9.0 Hz), 4.72 (d, 1H, J = 14 Hz), 4.64 (d, 1H, J = 14 Hz), 4.38-4.24 (m, 2H), 3.98-3.86 (m, 1H), 2.92 (d, 1H, J = 17 Hz), 2.65 (dd, 1H, J = 17, 12 Hz), 1.63 (s, 9H), 1.61 (s, 9H).

MS: 598 (M-1).

To a solution of 50 % trifluoroacetic acid/dichloromethane (1 ml) at room temperature was added oxalic acid 3-(*tert*-butoxycarbonyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H- thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (50 mg, 0.092 mmol). The solution was stirred for 3 hours.

The solvent was removed in vacuo. The residue was washed with dichloromethane affording after filtration 25 mg (73 %) of the title compound as a solid.

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 12.32 (s, 1H), 4.82 (d, 1H, J = 15 Hz), 4.68 (d, 1H, J = 15 Hz), 4.37 (s, 1H), 3.92 (m, 1H), 2.93 (d, 1H, J = 16 Hz), 2.60 (dd, 1H, J = 30, 10 Hz).

MS: 372 (M-1).

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# **EXAMPLE 22**

7-Hydroxymethyl-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a mixture of 2-hydroxymethyl-tetrahydro-pyran-4-one (35 g, 0.27 mol), tert-butyl cyanoacetate (58.68 ml g, 0.4 mol), and sulphur (9.47 g, 0.3 mol) in absolute ethanol (400 ml) was added morpholin (47 ml, 0.54 mol), and the resulting mixture was heated to 45 °C for 16 h. The reaction mixture was cooled, filtered and the filtrate evaporated in vacuo. The resultant oil was dissolved in ethyl acetate (600 ml), washed with water (3 x 200 ml), brine (200 m), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was crystallised from diethyl ether (100 ml) followed by addition of a mixture of diethyl ether and heptane (100 ml, 1:1). The precipitate was filtered off, washed with a mixture of diethyl ether and heptane (90 ml, 1:1) and dried in vacuo at 50 °C for 52 h affording 44.51 g of a mixture of 5 and 7 regioisomers according to NMR. The mixture of regioisomers (44.51 g) was suspended in diethyl ether (500 ml) and stirred

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at room temperature for 96 h. and at reflux temperature for 2 h. After cooling to room temperature the precipitate was filtered off and washed with a mixture of diethyl ether and heptane (100 ml, 1:1) which afforded after drying in vacuo at 50 °C, 22.12 g (29 %) of 2-amino-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

All filtrates were pooled and evaporated in vacuo affording 55 g of a mixture of regioisomers. To 40.16 g (0.141 mol) of this regioisomer mixture dissolved in dichloromethane (450 ml) was added diisopropylethylamine (49.5 ml, 0.28 mol) and the mixture was cooled to 0 °C. Chlorothiethylsilane (38.2 ml, 0.23 mol) was added dropwise and the mixture was stirred for 10 minutes and for 15 minutes at room temperature. The reaction mixture was washed with saturated aqueous sodium carbonate (3 x 150 ml), brine (3 x 150 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue (70.4 g) was partitioned into two portions which were subjected to flash chromatography (2 I silicagel) using a mixture of ethyl acetate/hexane (1:20) as eluant. Pure fractions of 2-amino-5-triethylsilanyloxymethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester and 2-amino-7-triethylsilanylhydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tertbutyl ester were collected. A fraction containing both isomers (18.84 g) was re-subjected to flash chromatography (2 I silicagel) using a mixture of ethyl acetate/hexane (1:20) as eluant. A total of 28.1 g (50 %) of 2-amino-5-triethylsilanylhydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester was obtained. A total of 18.2 g (32 %) of 2amino-7-triethylsilanylhydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester was obtained.

To the above 2-amino-7-triethylsilanylhydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (18.2 g, 0.046 mol) dissolved in dichloromethane (200-ml) was added a mixture of imidazol-1 yl-oxo-acetic acid *tert* butyl ester (17.9 g, 0.091 mol) in dichloromethane (30 ml) under nitrogen. The reaction mixture was allowed to stir at room temperature for 18 h. The reaction mixture was evaporated <u>in vacuo</u> and

the residue was dissolved in ethyl acetate (100 ml) and washed with 1 N hydrochloric acid (3 x 50 ml), brine (3 x 75 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the organic phase evaporated in vacuo affording in quantitative yield 2-(tert-butoxyoxalyl-amino)-7-triethylsilanyloxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester ester as a oil.

To a mixture of the above 7-triethylsilanyl ether (24.0 g, 0.046 mol) in tetrahydrofuran (100 ml) was added 1 N hydrochloric acid (18 ml) and the reaction mixture was stirred at room temperature for 1.5 h. Ethyl acetate (150 ml) was added and the reaction mixture was washed with saturated aqueous sodium carbonate (3 x 100 ml), brine (3 x 100 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was tritituated with a mixture of diethyl ether and heptane (1:5) and the precipitate was filtered off, washed with heptane and dried in vacuo at 50 °C for 16 h affording 13.55 g (57 %) of 2-(tert-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

The above 2-(*tert*-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (16 mg, 0.033 mmol) was dissolved in 50 % trifluoroacetic acid in dichloromethane (1 ml). The reaction was stirred at room temperature for 3 hours. The volatiles were evaporated in vacuo and the residue washed with dichloromethane which afforded 7 mg (73 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 12.32 (s, 1H), 4.62 (s, 1H), 4.12 (m, 1H), 3.62-3.78 (m, 2H), 3.40-3.52 (m, 1H), 2.83 (m, 2H).

MS: 300 (M-1).

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7-(2,4-Dioxo-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

- To a solution of 2-amino-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.13 g, 0.46 mmol) in tetrahydrofuran (3 ml) was added triphenylphosphine (0.13 g, 0.51 mmol), and 2,4-thiazolidinedione (60 mg, 0.51 mmol). The reaction mixture was cooled to 0 °C and diisopropylazodicarboxylate (99 μl,
- 10 0.51 mmol) was added via syringe. The resultant mixture was stirred for 18 hours, gradually warming to room temperature. The volatiles were evaporated in vacuo and the resulting oil was diluted in ethyl acetate (50 ml). The organic phase was washed with saturated sodium bicarbonate (3 x 50 ml), brine (3 x 50 ml), dried(MgSO<sub>4</sub>),
- filtered and the solvent evaporated <u>in vacuo</u>. The residue was subjected to flash chromatography using a mixture of dichloromethane/methanol (9:1) as eluant. Pure fractions were collected (R<sub>f</sub>=0.70) and the solvent evaporated <u>in vacuo</u> which afforded 89 mg (51 %) of 2-amino-7-(2,4-dioxo-thiazolidin-3-ylmethyl)-
- 20 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.
  - <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 6.02 (s, 2H), 4.82 (dm, 1H), 4.13-4.02 (bm, 2H), 3.99 (s, 2H), 3.75-3.67 (m, 1H), 3.60 (dd, 1H, J = 14, 3.3,), 2.81-2.74 (m, 2H), 1.54 (s, 9H).
- 25 MS: APCI (+): 385.6 (M+H).

To-a-solution of the above of 2-amino-7-(2,4-dioxo-thiazolidin-3-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (89 mg) in tetrahydrofuran (5 ml) was added imidazol-1-yl-

oxo-acetic acid *tert*-butyl ester (79 mg, 0.312 mmol) and the mixture allowed to stir overnight at room temperature. The volatiles were evaporated in vacuo, the residue diluted with ethyl acetate and subjected to preparative chromatography using a mixture of

- dichloromethane/methanol (9:1) as eluant. Material eluting with R<sub>f</sub>= 0.72 was collected and the solvent evaporated <u>in vacuo</u> affording 40 mg (25 %) of 2-(*tert*-butoxyoxalyl-amino)-7-(2,4-dioxo-thiazolidin-3-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.
- <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.53 (s, 1H), 5.03 (dm, 1H), 4.12-4.04 (m, 2H), 4.01 (s, 2H), 3.79-3.71 (m, 2H), 2.88 (m, 2H), 1.62 (s, 9H), 1.59 (s, 9H).
   MS: APCI (+): 513.3 (M+H).
- The above 2-(*tert*-butoxyoxalyl-amino)-7-(2,4-dioxo-thiazolidin-3-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (40 mg) was dissolved in 50 % trifluoroacetic acid in dichloromethane (1 ml) and stirred at room temperature for 3 hours. The mixture was concentrated <u>in vacuo</u>, the residue titurated with
- dichloromethane and methanol which afforded after drying in vacuo 18 mg (87 %) of the title compound as a solid.
   <sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub> + CD<sub>3</sub>OD) δ 4.98 (dm, 1H), 4.16 (s, 2H), 4.14-4.02 (m, 2H), 3.78-3.72 (m, 2H), 2.91 (m, 2H).
   APCI (-): 399 (M-H);
- 25 LC-MS: s, 99%.

# **EXAMPLE 24**

# 7-(1,3-Dioxo-1,3-dihydro-isoindol-2-yloxymethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a mixture of 2-(tert-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (0.5 g, 1.2 mmol), 2-hydroxy-isoindole-1,3-dione (0.21 g, 1.3 mmol) and triphenylphosphine (0.35 g, 1.33 mmol) in dry tetrahydrofuran (20 ml) cooled to 0 °C under a nitrogen atmosphere was added diethyl azodicarboxylate (DEAD) (205 µl, 1.33 mmol). The reaction mixture was allowed to stir overnight, slowly warming to room temperature. The volatiles were evaporated in vacuo and the resultant solid dissolved in ethyl acetate (50 ml). The organic phase was washed with saturated aqueous sodium hydrogencarbonate (3 x 30 ml), water (3 x 50 ml), dried(Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated in vacuo. The residue (1.02 g) was subjected to flash column chromatography (300 ml silicagel) using a mixture of ethyl acetate/hexane (1:2) as eluant. Pure fractions were collected affording after evaporation in vacuo 0.37 g (54 %) of 2-(tert-butoxyoxalyl-amino)-7-(1,3-dioxo-1,3-dihydro-isoindol-2vloxymethyl)-4.7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil.

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The above di-*tert*-butyl ester (0.33 g, 0.59 mmol) was dissolved in 25 % trifluoroacetic acid in dichloromethane (2 ml). The reaction was stirred at room temperature for 6.5 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with a mixture of diethyl ether and heptane (5 ml, 1:1). The precipitate was filtered off, washed with heptane and diethyl ether, dried <u>in vacuo</u> at 50 °C for 18 h which afforded 200 mg (77 %) of the title compound as a solid.

30 M.p.: 251.5 - 254 °C;

Calculated for C<sub>19</sub>H<sub>14</sub>N<sub>2</sub>O<sub>9</sub>S;

C, 51.12 %; H, 3.16 %; N, 6.28 %. Found:

C, 51.46 %; H, 3.71 %; N, 5.87 %.

### EXAMPLE 25

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7-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 4-hydroxy-isobenzofuran-1,3-dione (0.5 g, 3.03 mmol) in anhydrous N,N-dimethylformamide (6 ml) under nitrogen was added diisopropylethylamine (1.05 ml, 6.06 mmol). The solution was stirred with cooling in an ice bath and chloromethyl methyl ether (0.46 ml, 6.06 mmol) was added. The reaction was allowed to slowly warm to ambient temperature and then stirred for an additional 7 h. The mixture was concentrated in vacuo to a small volume and diluted with ethyl acetate (75 ml). The organic layer was washed with water (2 x 40 ml), brine (20 ml), dried(Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo to give 0.6 g (95 %) of 4-methoxymethoxy-isobenzofuran-1,3-dione as a solid.  $^{1}$ H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.81 (t, J = 8, 1H), 7.62 (d, J = 8, 1H), 7.59 (d, J = 8, 1H), 5.43 (s, 2H), 3.55 (s, 3H).

A mixture of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.15 g, 0.53 mmol) and 4-methoxymethoxy-isobenzofuran-1,3-dione (135 mg, 0.64 mmol) was dissolved in distilled acetonitrile (7 ml) under nitrogen. The flask was cooled in an ice bath with stirring and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.12 g, 0.64 mmol), and triethylamine (0.22 ml, 1.59 mmol) were added. The reaction was warmed to ambient temperature and stirred for 18 h. The solution was concentrated <u>in vacuo</u>

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and the residue dissolved in ethyl acetate (40 ml). The organic layer was washed with 1% hydrochloric acid (2 x 10 ml), saturated sodium bicarbonate (10 ml), and brine (10 ml). The resulting solution was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo which afforded 0.18 g of a crude 2-amino-7-(4-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester which was used without further purification.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.65-7.58 (m, 2H), 7.51 (d, *J* = 8, 1H), 6.00-5.86 (2s, 2H), 5.39 (s, 2H), 4.94-4.89 (m, 1H), 4.18-4.02 (m, 2H), 3.86-3.65 (m, 2H), 3.54 (s, 3H), 2.85-2.73 (m, 2H), 1.55 (s, 9H). APCI-MS: [M+H]<sup>+</sup> = 475.4

To a solution of crude 2-amino-7-(4-methoxymethoxy-1,3-dioxo-1,3dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester (0.18 g) in distilled dichloromethane (4 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid tert-butyl ester (0.23 g, 1.2 mmol). The reaction was stirred for 3 hours., concentrated in vacuo and reconstituted in ethyl acetate (30 ml). The organic layer was washed with 1% hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The resulting solution was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (0 to 5 % gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo to give 90 mg (28 % in two steps) of 2-(tert-butoxyoxalyl-amino)-7-(4-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 12.54 (s, 1H), 7.64 (t, J = 8, 1H), 7.51 (d, J = 8) 8, 1H), 7.46 (d, J = 8, 1H), 5.40 (s, 2H), 5.11-5.07 (m, 1H), 4.16-4.08 (m, 2H), 3.84-3.72 (m, 2H), 3.55 (s, 3H), 2.95-2.81 (m, 2H), 1.62 (s, 9H), 1.59 (s, 9H). ...

APCI-MS:  $[M+H]^{+} = 603.8$ 

The above 2-(*tert*-butoxyoxalyl-amino)-7-(4-methoxymethoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (86 mg, 0.143 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (4 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 55 mg (86 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (400 MHz, d<sub>6</sub>-DMSO) δ 12.34 (s, 1H), 11.10 (s, 1H), 7.63 (t, J = 8, 1H), 7.31 (d, J = 8, 1H), 7.22 (d, J = 8, 1H), 4.99-4.95 (m, 1H), 4.05-4.00 (m, 1H), 3.91-3.86 (m, 1H), 3.76-3.66 (m, 2H), 2.88-2.80 (m, 2H). APCI-MS: [M+H]<sup>†</sup> = 447.4

HPLC (254.4nm): R<sub>t</sub>=2.921 min, 100%

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### **EXAMPLE 26**

25 <u>7-(5-Methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid</u>

The <u>title compound</u> was prepared in a similar way as described in Example 25.

M.p.: 234 - 236 °C;

Calculated for  $C_{20}H_{16}N_2O_9S$ , 0.25 x  $H_2O$ ;

C, 51.67 %; H, 3.58 %; N, 6.03 %. Found:

5 C, 51.95 %; H, 3.92 %; N, 6.06 %.

## **EXAMPLE 27**

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7-(5,7-Dioxo-5,7-dihydro-[1,3]dioxolo[4,5-f]isoindol-6-ylmethyl2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in 15 Example 25.

M.p.: 239.5 - 242.5 °C;

Calculated for  $C_{20}H_{14}N_2O_{10}S$ , 0.1 x  $H_2O$ ;

C, 50.45 %; H, 3.01 %; N, 5.88 %. Found:

20 C, 51.06 %, H, 3.43 %; N, 5.93 %.

# **EXAMPLE 28**

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# 7-(((Benzo[1,3]dioxole-5-carbonyl)-amino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

Phthalimidoacetaldehyde diethyl acetal (100 g, 0.38 mol) and 1 N 5 hydrochloric acid (600 ml) was mixture was stirred at reflux temperature for 5 min. or until a homogeneous solution is obtained. The reaction mixture was cooled and the precipitate was filtered off and dried in vacuo at 50 °C for 16 h which afforded 63.3 g (88 %) of phthalimidoacetaldehyde as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 4.58 (s, 2H), 7.76 - 7.78(m, 2H), 7.90 - 7.92 (m, 2H), 9.67 (s, 1H).

To a mixture of phthalimidoacetaldehyde (64 g, 0.34 mol) and trans-1methoxy-3-(trimethylsilyloxy)-1,3-butadiene (81.5 g, 0.38 mol) in benzene (600 ml) stirred for 15 min. under nitrogen was added dropwise a 45 % solution of zinc chloride diethyl ether complex in dichloromethane (55.5 ml, 0.17 mol) at 0 °C. The reaction was allowed warm up to room temperature overnight. To the reaction mixture was added water (500 ml) and the resulting mixture was extracted with ethyl acetate (200 ml). The organic extract was washed successively with 1.0 N hydrochloric acid (2 x 200 ml) and brine (200 ml). The organic phase was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo which afforded a slowly crystallising oil (98 g). To the solid was added a mixture of ethyl acetate and diethyl ether (400 ml, 1:1) and the resulting precipitate was filtered off, washed with a small portion of diethyl ether and dried at 50 °C for 1h affording 59.8 g (69 %) of 2-(4-oxo-3,4-dihydro-2H-pyran-2-ylmethyl)isoindole-1,3-dione as a solid. The filtrate was evaporated in vacuo and the residue purified by column chromatography on silica gel (1 L) using a mixture of ethyl acetate and heptane (1:2) as eluant. Pure fractions were collected and the solvent evaporated in vacuo to almost dryness, the solid was filtered off and dried in vacuo-at-50 °C for 16 h affording an additional 15 g (17 %) of 2-(4-oxo-3,4-dihydro-2H-pyran-2-ylmethyl)-isoindole-1,3dione as a solid.

 $^{1}$ H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  2.61 (d, 2H), 3.85 (dd, 1H), 4.18 (dd, 1H), 4.76 (m, 1H), 5.43 (d, 1H), 7.28 (d, 1H), 7.69 - 7.77 (m, 2H), 7.84 - 7.88 (m, 2H).

2-(4-Oxo-3,4-dihydro-2H-pyran-2-ylmethyl)-isoindole-1,3-dione (13 g, 5 0.051 mol) was dissolved in ethyl acetate (250 ml) and placed in a Parr bottle. 10 % Pd/C (1.5 g) was carefully added and the mixture was shaken under a pressure of 30 psi of hydrogen for 6.5 h (Parr apparatus). Filtration followed by evaporation of the ethyl acetate in vacuo afforded a crude 11.5 g of 2-(4-oxo-tetrahydro-pyran-2-ylmethyl)-isoindole-1,3-dione 10 pure enough for the next step. Analytical pure compound could be obtained by purification of a small sample (250 mg) by column chromatography on silica gel, utilising a mixture of hexane/ethyl acetate as a gradient (from 100/0 to 50/50). Pure fractions were collected and the solvent evaporated in vacuo affording 142 mg (55 %) of 2-(4-oxo-15 tetrahydro-pyran-2-ylmethyl)-isoindole-1,3-dione as a solid. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  2.30 - 2.68 (m, 4H), 3.62 (m, 1H), 3.74 (m, 1H), 4.00 (m, 2H), 7.75 (m, 2H), 7.88 (m, 2H).

To a mixture of 2-(4-oxo-tetrahydro-pyran-2-ylmethyl)-isoindole-1,3-dione 20 (11.5 g, 44 mmol), tert-butyl cyanoacetate (6.9 g, 49 mmol) and elemental sulfur (1.6 g, 49 mmol) in ethanol (250 ml) was added morpholin (15 ml) and the resulting mixture was stirred at 50 °C for 16 h. The cooled reaction mixture was filtered and the precipitate filtered off and washed with diethyl ether and dried in vacuo affording 6.5 g (35 %) of 2-amino-5-(1,3-dioxo-25 1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3carboxylic acid tert-butyl ester as a solid. The filtrate was evaporated in vacuo and the residue was dissolved in ethyl acetate (200 ml) washed with water (2 x 100 ml), brine (100 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo affording 6.0 g-(33-%) of almost regioisomer-pure 2-amino-7-(1,3-dioxo-1,3-dihydroisoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 1.50 (s, 9H), 2.54 - 2.63 (m, 1H), 2.84 - 2.90 (m, 1H), 3.79 (q, 1H), 3.96 - 4.04 (m, 2H), 4.48 - 4.62 (m, 2H), 5.91 (bs, 2H, N*H*<sub>2</sub>), 7.70 (m, 2H), 7.84 (m, 2H).

To a solution of 2-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (6.0 g, 0.014 mol) in ethanol (100 ml) was added hydrazine-hydrate (1.4 ml, 0.029 mol). The mixture was stirred at reflux temperature for 1 h. The cooled reaction mixture was filtered and the solvent evaporated <u>in vacuo</u>. The residue was dissolved in diethyl ether (200 ml) and washed with water (100 ml), brine (100 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u> affording 2.9 g (71 %) of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

To a ice cooled mixture of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (1.4 g, 4.92 mmol), triethylamine (2 ml) in dichloromethane (100 ml) was added dropwise a solution of benzo[1,3]dioxole-5-carbonyl chloride (1.0 g, 5.41 mmol) in dichloromethane (25 ml) during 1.5 h. The ice cooled reaction mixture was stirred for an additional 0.5 h. The volatiles were evaporated <u>in vacuo</u> and the residue was dissolved in ethyl acetate (200 ml) and washed with water (2-x-100 ml), brine (100-ml), dried-(Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue (2 g) was subjected to flash column chromatography (1 I silicagel) using a mixture of ethyl acetate/hexane

(1:2) as eluant. Pure fractions were collected affording after evaporation in vacuo 0.3 g (14 %) of 2-amino-7-(((benzo[1,3]dioxole-5-carbonyl)amino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

5 TLC:  $R_f = 0.44$  (ethyl acetate/heptane 1:1)

A mixture of the above 2-amino-7-(((benzo[1,3]dioxole-5-carbonyl)amino)methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.3 g, 0.69 mmol), imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.16 g, 0.83 mmol) in dry tetrahydrofuran (50 ml) was stirred at room temperature for 16 h. The volatiles were evaporated in vacuo and the residue was dissolved in ethyl acetate (100 ml) and washed with water (2 x 50 ml), brine (50 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue (0.35 g) was subjected to flash column chromatography (500 ml silicagel) using a mixture of ethyl acetate/hexane (1:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo. The residue was trituated with diethyl ether (5 ml), filtered off and dried in vacuo at 50 °C for 5 h which afforded 0.17 g (44 %) of 7-(((benzo[1,3]dioxole-5-carbonyl)amino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

TLC:  $R_f = 0.37$  (ethyl acetate/heptane 1.1).

The above di-*tert*-butyl ester ( 0.17 g, 0.30 mmol) was dissolved in 25 % trifluoroacetic acid in dichloromethane (20 ml). The reaction was stirred at room temperature for 5.5 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with diethyl ether (10 ml). The precipitate was filtered off, washed with diethyl ether, dried <u>in vacuo</u> at 50 °C for 72 h which afforded 100 mg (74 %) of the <u>title compound</u> as a solid.

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M.p.: 227 - 230° C;

Calculated for  $C_{19}H_{16}N_2O_9S$ , 0.5 x  $H_2O$ ;

C, 49.89 %; H, 3.75 %; N, 6.12 %. Found:

C, 50.02 %; H, 3.68 %; N, 5.98 %.

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### EXAMPLE 29

7-[3-(2,4-Dimethoxy-phenyl)-ureidomethyl]-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester (64 mg, 0.22 mmol) in dichloromethane (1 ml) was added 2,4-dimethoxyphenylisocyanate (40 mg, 0.22 mmol). The reaction mixture was stirred at room temperature for 18 h. The reaction mixture was concentrated in vacuo, diluted with ethyl acetate (30 ml), washed with saturated sodium carbonate (3 x 25 ml), brine (3 x 25 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was subjected to preparative thin layer chromatography (100% dichloromethane). R<sub>f</sub>=0.8 was isolated and the solvent evaporated in vacuo which afforded 55 mg (53 %) of 2-amino-7-(3-(2,4-dimethoxyphenyl)ureidomethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  7.75 (d, J = 9.6, 1H), 7.62 (d, J = 8.1, 1H), 6.45 (m, 3H), 5.00 (bs, 2H), 4.68 (m, 1H), 4.12 (m, 2H), 3.80 (s, 3H), 3.76 (s, 3H), 3.76-3.67 (m, 1H), 3.30 (dd, J = 14, 6.9, 1H), 2.76

(m, 2H), 1.55 (s, 9H).

MS: APCI (+): 464.3 (M+H).

To a solution of the above 2-amino-7-(3-(2,4-dimethoxyphenyl)ureidomethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (47 mg, 0.11 mmol) in dichloromethane (1 ml) was

added triethylamine (28 µl, 0.22 mmol) and midazol-1-yl-oxo-acetic acid tert-butyl ester (40 mg, 0.22 mmol). The mixture allowed to stir at room temperature for 18 h. The volatiles were evaporated in vacuo and the residue diluted with ethyl acetate (35 ml). The organic phase was washed with saturated sodium carbonate (3 x 25 ml), brine (3 x 25 ml), dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The resultant oil was subjected to preparative thin layer chromatography (60 % ethyl acetate/40 % hexanes). Pure 2-(tertbutoxyoxalyl-amino)-7-(3-(2,4-dimethoxy-phenyl)ureidomethyl)-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester 34 mg 10 (58 %) was isolated as an oil. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  12.49 (s, 1H), 7.70 (d, J = 9.6, 1H), 6.62 (bs, 1H), 6.47 (m, 3H), 5.02 (bs, 1H), 4.84 (m, 1H), 4.19 (dm, 1H), 3.82 (s, 3H), 3.79 (s, 3H), 3.75-3.70 (m, 1H), 3.36 (dd, J = 13.5, 7.5, 3.751H), 2.87 (m, 2H), 1.61 (s, 9H), 1.60 (s, 9H). 15 MS: APCI (+): 592.4 (M+H).

The above 2-(*tert*-butoxyoxalyl-amino)-7-(3-(2,4-dimethoxy-phenyl)ureidomethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (34 mg) was dissolved in 20 % trifluoroacetic acid in dichloromethane (2 ml) and stirred at room temperature for 3 hours. The volatiles were evaporated <u>in vacuo</u> and the residue was titurated with diethyl ether (2x), filtered off and washed with a small amount of dichloromethane which afforded after drying <u>in vacuo</u> 16 mg (89 %) of the title compound as a solid.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ 7.66 (d, J = 9, 1H), 6.53 (d, J = 2.7, 1H), 6.44 (dd, J = 9, 2.7, 2H), 4.82 (m, 1H), 4.2 (m, 2H), 3.82 (s, 3H), 3.76 (s, 3H), 3.67 (dd, J = 13, 4.5, 2H), 2.94 (m, 2H). MS: APCI (+): 480.3 (M+H);

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#### **EXAMPLE 30**

2-(Oxalyl-amino)-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

A solution of glyoxylic acid ethyl ester, polymer form (2.02 g, 8.9 mmol) and (3-methoxy-1-methylene-allyloxy)-trimethyl-silane (1.9 ml, 8.9 mmol, Danishefsky's diene) in benzene (12 ml) was placed under nitrogen. Zinc chloride (0.5N in tetrahydrofuran, 8.9 ml, 4.45 mmol) was added and the reaction stirred at ambient temperature for 72 h. The mixture was concentrated in vacuo, diluted with ethyl acetate (100 ml) and washed with 1N hydrochloric acid (20 ml), saturated sodium bicarbonate (20 ml), and brine (20 ml). The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a mixture of ethyl acetate/hexane (1:2) as eluant. Pure fractions were collected and the solvent evaporated in vacuo which afforded 1.2 g (75 %) of 4-oxo-3,4-dihydro-2H-pyran-2-carboxylic acid ethyl ester as an oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.40 (d, J = 6, 1H), 5.48 (d, J = 6, 1H), 5.01 (t, J = 8, 1H), 4.28 (q, J = 7, 2H), 2.85 (d, J = 8, 2H), 1.29 (t, J = 7, 3H).

To a solution of the above of 4-oxo-3,4-dihydro-2H-pyran-2-carboxylic acid ethyl ester (1.0 g, 5.9 mmol) in ethyl acetate (12 ml) was added 10 % palladium on activated carbon (0.15 g). The reaction was shaken on a Parr hydrogenator under a hydrogen atmosphere (30 psi) for 1.5 h. The mixture was filtered through celite and concentrated in vacuo. The residue was purified by silica gel chromatography using diethyl ether as eluant. Pure-fractions-were-collected and the solvent evaporated in vacuo which affording 0.6 g (60 %) of 4-oxo-tetrahydro-2H-pyran-2-carboxylic acid ethyl as an oil.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  4.41-4.35 (m, 1H), 4.26 (q, J = 7, 2H), 3.81-3.70 (m, 1H), 2.73-2.58 (m, 3H), 2.44-2.36 (m, 1H), 1.29 (t, J = 7, 3H).

To a solution of 4-oxo-tetrahydro-2H-pyran-2-carboxylic acid ethyl (0.6 g, 3.5 mmol) in absolute ethanol (6 ml) was added sulfur (0.12 g, 3.85 mmol) and tert-butyl cyanoacetate (0.64 g, 4.55 mmol). The solution was stirred under nitrogen in a 50 °C oil bath and morpholin (0.61 ml, 7.0 mmol) was added. The reaction was stirred for 18 h. and then cooled to ambient temperature and excess sulfur removed by filtration. The filtrate was concentrated in vacuo and reconstituted in ethyl acetate (50 ml). The organic phase was washed with brine (2 x 10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/hexane (20 to 25 % gradient) as eluant. Pure fraction of the two isomers were collected and the solvent evaporated in vacuo which afforded 0.47 g of 2-amino-4,7dihydro-5H-thieno[2,3-c]pyran-3,5-dicarboxylic acid 3-tert-butyl ester 5ethyl ester (A) and 0.3 g of 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 3-tert-butyl ester 7-ethyl ester (B) in 62 % combined yield.

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(A)

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  5.96 (bs, 2H), 4.77-4.61 (m, 2H), 4.32-4.18 (m, 3H), 3.19-3.12 (m, 1H), 2.90-2.80 (m, 1H), 1.52 (s, 9H), 1.29 (t, J = 7, 3H).

25 APCI-MS:  $[M+H]^+ = 272.4$  (loss of t-butyl)

(B)

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  5.10 (s, 1H), 4.28-4.13 (m, 3H), 3.98-3.91 (m, 1H), 2.82-2.76 (m, 2H), 1.51 (s, 9H), 1.31 (t, J = 7, 3H).

30 APCI-MS: [M+H]<sup>+</sup> = 272.4 (loss of t-butyl)

The above 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,5-dicarboxylic

acid 3-tert-butyl ester 5-ethyl ester (275 mg, 0.84 mmol) was dissolved in

a mixture of ethanol (4 ml) and tetrahydrofuran (1 ml). Sodium hydroxide

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(1N, 1.6 ml, 1.68 mmol) was added and the reaction stirred at ambient temperature for 5 h. after which TLC analysis indicated that the reaction was complete. The reaction was monitored with a pH meter and neutralized with 1N hydrochloric acid until pH = 6.9. The solution was concentrated in vacuo to give 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,5-dicarboxylic acid 3-tert-butyl ester as a solid. Sodium chloride remained as an impurity.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ 4.67-4.54 (m, 2H), 4.00-3.95 (m, 1H), 3.20-3.12 (m, 1H), 2.74-2.63 (m, 1H), 1.54 (s, 9H).

10 APCI-MS:  $[M+H]^+ = 300.0$ 

To a solution of the above 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,5-dicarboxylic acid 3-*tert*-butyl ester (94 mg, 0.31 mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (72 mg, 0.37 mmol) in distilled dichloromethane (4 ml) under nitrogen was added aniline (32  $\mu$ l, 0.34 mmol) followed by 2,6-lutidine (0.11 ml, 0.93 mmol). The reaction was stirred for 72 h., concentrated in vacuo and reconstituted in ethyl acetate (30 ml). The organic layer was washed with 1% hydrochloric acid (10 ml), saturated sodium bicarbonate (10 ml), brine (10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo to give 51 mg (45 %) of 2-amino-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  8.40 (s, 1H), 7.60 (d, 1H, J = 7), 7.49 (d, 1H, J = 8), 7.34 (t, 1H, J = 8), 7.32 (t, 1H, J = 8), 7.13 (t, 1H, J = 7), 6.03 (s, 2H), 4.82-4.73 (m, 2H), 4.25-4.22 (m, 1H), 3.43-3.38 (m, 1H), 2.79-2.72

APCI-MS:  $[M+H]^{+} = 375.5$ 

(m, 1H), 1.54 (s, 9H).

To a solution of the above 2-amino-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (51 mg, 0.14 mmol) in distilled dichloromethane-(3-ml)-under\_nitrogen was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (80 mg, 0.42 mmol) and triethylamine (38 μl, 0.28 mmol). The reaction was stirred for 4 h., concentrated <u>in vacuo</u>

and reconstituted in ethyl acetate (25 ml). The organic layer was washed with 1% hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), brine (5 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated <u>in vacuo</u>. The crude material was purified by silica gel chromatography using a 4 % mixture of ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> to give 41 mg (26 % over two steps) of 2-(*tert*-butoxyoxalyl-amino)-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.56 (s, 1H), 8.40 (s, 1H), 7.59 (d, J = 8, 2H), 7.33 (t, J = 8, 2H), 7.12 (t, J = 7, 1H), 5.01-4.85 (m, 2H), 4.27-4.22 (m, 1H), 3.54-3.47 (m, 1H), 3.89-2.79 (m, 1H), 1.60 (s, 9H), 1.58 (s, 9H). *APCI-MS*: [M+H]<sup>+</sup> = 503.2

The above 2-(*tert*-butoxyoxalyl-amino)-5-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (37 mg, 0.074 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (3 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with ethyl ether and dried <u>in vacuo</u> to give 18 mg (62 %) of the <u>title compound</u>.

1 H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.32 (s, 1H), 9.85 (s, 1H), 7.69 (d, *J* = 8, 2H), 7.31 (t, *J* = 8, 2H), 7.07 (t, *J* = 7, 1H), 4.98 (d, *J* = 15, 1H), 4.83 (d, *J* = 15, 1H), 4.35-4.31 (m, 1H), 3.23 (d, *J* = 17, 1H), 2.84 (dd, *J* = 17, 10, 1H).

25 APCI-MS:  $[M+H]^{+} = 391.3$ HPLC (254.4nm):  $R_{t}=3.22$  min, 100%

#### **EXAMPLE 31**

# 5-Benzylcarbamoyl-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,5-dicarboxylic acid 3-*tert*-butyl ester (101 mg, 0.34 mmol, prepared in Example 31) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (78 mg, 0.41 mmol) in distilled dichloromethane (4 ml) under nitrogen was added benzylamine (40 μl, 0.37 mmol) followed by 2,6-lutidine (0.12 ml, 1.02 mmol). The reaction was stirred for 72 h., concentrated <u>in vacuo</u> and reconstituted in ethyl acetate (30 ml). The organic layer was washed with 1 % hydrochloric acid (10 ml), saturated sodium bicarbonate (10 ml), brine (10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>) over sodium sulfate, filtered, and the solvent evaporated <u>in vacuo</u> to give 72 mg (56 %) of 2-amino-5-benzylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.36-7.28 (m, 5H), 4.66 (s, 2H), 4.44 (d, J = 5, 2H), 4.17-4.13 (m, 1H), 3.40-3.33 (m, 1H), 2.75-2.66 (m, 1H), 1.54 (s, 9H).

APCI-MS:  $[M+H]^{+} = 389.5$ 

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To a solution of the above 2-amino-5-benzylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3,carboxylic acid *tert*-butyl ester (72 mg, 0.19 mmol) in distilled dichloromethane (4 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.11 g, 0.57 mmol) and triethylamine (51 μl, 0.38 mmol). The reaction was stirred for 4 h., concentrated in vacuo and reconstituted in ethyl acetate (25 ml). The organic layer was washed with 1% hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), brine (5 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The crude material was purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (5 to 10 % gradient) as eluant. Pure fractions were collected and the solvent-evaporated-in-vacuo to give 42 mg (24 % over two steps) of 5-benzylcarbamoyl-2-(*tert*-butoxyoxalyl-

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amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 12.56 (s, 1H), 7.37-7.29 (m, 5H), 6.97 (t, 1H, J = 6), 4.89-4.77 (m, 2H), 4.58-4.46 (m, 2H), 4.20-4.16 (m, 1H), 3.50-3.44 (m, 1H), 2.84-2.76 (m, 1H), 1.61 (s, 9H), 1.60 (s, 9H). APCI-MS: [M+H]<sup>+</sup> = 517.3

The above 5-benzylcarbamoyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,carboxylic acid *tert*-butyl ester (36 mg, 0.07 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (3 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 14 mg (50 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 8.48 (t, J = 6, 1H), 7.31-7.20 (m, 5H), 4.91 (d, J = 15, 1H), 4.76 (d, J = 15, 1H), 4.32-4.29 (m, 2H), 4.20-4.16 (m, 1H), 3.22 (m, 1H, partially obscured by water), 2.70-2.63 (m, 1H).

APCI-MS:  $[M+H]^+ = 405.2$ 

20 ... HPLC (254.4nm): R<sub>t</sub>=3.06 min, 100 %

#### **EXAMPLE 32**

25 <u>2-(Oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid</u> 7-ethyl ester

To a solution of 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 3-tert-butyl ester 7-ethyl ester (60 mg, 0.18 mmol) in distilled dichloromethane (3 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid tert-butyl ester (0.11 g, 0.54 mmol) and triethylamine (50 µl, 0.36 mmol). The reaction was stirred for 4 h., concentrated in vacuo

and reconstituted in ethyl acetate (20 ml). The organic layer was washed with 1 % hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), brine (5 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated <u>in vacuo</u>. The crude material was purified by silica gel chromatography using a 6 % mixture of ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 78 mg (95 %) of 2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 3-*tert*-butyl ester 7-ethyl ester as an oil.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.54 (s, 1H), 5.28 (s, 1H), 4.27 (q, 2H, J = 7), 4.25-4.18 (m, 1H), 4.04-3.96 (m, 1H), 2.96-2.80 (m, 2H), 1.60 (s, 9H), 1.57 (s, 9H).

LC-MS: R<sub>t</sub>=3.97 min,  $[M+H]^+$  = 456.3

The above 2-(tert-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 3-tert-butyl ester 7-ethyl ester (72 mg, 0.16 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (4 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and the residue evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 48 mg (88 %) of the <u>title</u>

<sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.34 (s, 1H), 5.47 (s, 1H), 4.19 (q, J = 7, 2H), 3.98-3.94 (m, 2H), 2.90-2.78 (m, 2H), 1.23 (t, J = 7, 3H).

APCI-MS:  $[M+H]^{+} = 344.2$ 

compound as a solid.

HPLC (254.4nm):  $R_t$ =2.82 min, 100 %

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#### EXAMPLE 33

7-Benzylcarbamoyl-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-

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To a solution of 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7dicarboxylic acid 3-tert-butyl ester 7-ethyl ester (0.12 g, 0.37 mmol) in ethanol (3 ml) was added potassium hydroxide (56 mg, 1.0 mmol) dissolved in a minimum amount of water. The mixture was stirred for 24 h., then 1N hydrochloric acid was added until pH = 7. The solution was concentrated in vacuo and the residue partitioned between ethyl acetate (35 ml) and water (10 ml). The layers were separated and 1 % hydrochloric acid (1 ml) was added to the aqueous layer. The aqueous layer was then extracted further with ethyl acetate (3 x 15 ml) and the combined organic extracts were washed with brine, dried (Na<sub>2</sub>SO<sub>4</sub>) and filtered. Triethylamine (3 drops) was added to the solution to stabilize the acid-sensitive compound. The solution was concentrated in vacuo affording 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-dicarboxylic acid 3-tert-butyl ester triethylamine salt (approximately 0.13 g) as a solid. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  5.01 (s, 1H), 4.28-4.23 (m, 1H), 3.90-3.85 (m. 1H), 2.88-2.71 (m, 3H), 1.56 (s, 9H). A solution of the above 2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3,7-

dicarboxylic acid 3-*tert*-butyl ester triethylamine salt (0.12 g, 0.30 mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (71 mg, 0.36 mmol) was prepared in distilled acetonitrile under nitrogen. Benzylamine (36  $\mu$ l, 0.33 mmol) was added followed by 2,6-lutidine (70  $\mu$ l, 0.60 mmol). The reaction was stirred at ambient temperature for 18 h., then concentrated <u>in vacuo</u> and reconstituted in ethyl acetate (30 ml). The organic layer was washed with 1 % hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (2 x 5 ml), and brine (10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated <u>in vacuo</u> which afforded crude 2-amino-7-benzylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester which was used without purification.

To a solution of the above crude 2-amino-7-benzylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (77 mg, 0.2 mmol) in distilled dichloromethane (3 ml) under nitrogen was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.11 g, 0.6 mmol) and triethylamine (55 µl,

0.4 mmol). The reaction was stirred for 5 h., concentrated <u>in vacuo</u> and reconstituted in ethyl acetate (20 ml). The organic layer was washed with 1 % hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (5 ml), brine (5 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated <u>in vacuo</u>.

- The crude material was purified by silica gel chromatography using a 5 % mixture of ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 29 mg (19 % over two steps) of 7-benzylcarbamoyl-2-(tert-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil.
- <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 12.49 (s, 1H), 7.35-7.26 (m, 5H), 6.96 (t, J = 6, 1H), 5.20 (s, 1H), 4.55-4.41 (m, 2H), 4.22-4.17 (m, 1H), 3.87-3.81 (m, 1H), 2.97-2.84 (m, 2H), 1.61 (s, 9H), 1.59 (s, 9H).

APCI-MS:  $[M-H]^{-} = 516$ 

The above 7-benzylcarbamoyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (29 mg, 0.06 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (2 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and the residue evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 18 mg (80 %) of the <u>title</u> compound as an solid.

<sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.33 (s, 1H), 8.67 (t, J = 6, 1H), 7.30-7.21 (m, 5H), 5.23 (s, 1H), 4.31-4.28 (m, 2H), 4.13-4.10 (m, 1H), 3.88-3.85 (m, 1H), 2.86 (bs, 2H).

25 APCI-MS:  $[M+H]^+ = 405$ HPLC (254.4nm):  $R_t=3.12$  min, 94 %

# EXAMPLE 34

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# 7-((2-(4-Methanesulfonyl-phenyl)-acetylamino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of (4-methanesulfonyl-phenyl)-acetic acid (90.4 mg, 0.42 mmol) in a mixture of dichloromethane (3 ml) and N,Ndimethylformamide (1 ml) cooled at 0 °C was added diisopropylethylamine (306 µl, 1.76 mmol), diisopropylazodicarboxylate (72 µl, 0.45 mmol) and 1-hydroxy-benzotriazole (56.6 mg, 0.42 mmol). After being stirred for 20 minutes, 2-amino-7-aminomethyl-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (100 mg, 0.35 10 mmol) dissolved in dichloromethane (1 ml) was added via syringe. The reaction mixture was stirred for 18 h. while slowly warming to room temperature. The volatiles were evaporated in vacuo and the residue diluted with ethyl acetate (50 ml). The organic phase was washed with saturated sodium bicarbonate (3 x 50 ml), 1 % 15 hydrochloric acid (3 x 50 ml), brine (3 x 50 ml), dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The resultant oil was subjected to preparative thin layer chromatography using a mixture of methanol/dichloromethane (1:9) as eluant. Fraction with R<sub>f</sub>=0.5 was isolated which afforded after evaporating the solvent in vacuo 115 mg 20 (69 %) of 2-amino-7-((2-(4-methanesulfonyl-phenyl)acetylamino)methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 7.87 (d, *J* = 8.7, 2H), 7.39 (d, *J* = 7.5, 2H), 5.91 (bs, 2H), 4.65 (m, 1H), 4.09 (dt, J = 7.8, 3.3, 1H), 3.85-3.65 (m, 2H), 3.61 (s, 2H), 3.45-3.38 (m, 2H), 3.05 (s, 3H), 2.75 (m, 2H), 1.56 (s, 9H).

MS: APCI (+): 481 (M+H).

To a solution of the above 2-amino-7-((2-(4-methanesulfonyl-phenyl)acetylamino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (110 mg, 0.23 mmol) in dichloromethane (3 ml) was added triethylamine (96 μl, 0.69 mmol) and midazol-1-yl-oxo-acetic acid *tert*-butyl ester (134 mg, 0.69 mmol).

The reaction was stirred at room temperature for 18 h. The reaction mixture was concentrated <u>in vacuo</u>, diluted in ethyl acetate (50 ml), washed with saturated sodium carbonate (3 x 50 ml), brine (3 x 50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The resultant oil was subjected to preparative thin layer chromatography using a mixture of methanol/dichloromethane (1:9). Fraction with R<sub>f</sub>=0.5 was collected and the solvent evaporated <u>in vacuo</u> affording 70 mg (50 %) of 2-(*tert*-butoxyoxalyl-amino)-7-((2-(4-methanesulfonyl-phenyl)acetylamino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-

- carboxylic acid *tert*-butyl ester as an oil. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  12.49 (s, 1H), 7.88 (d, J = 8.1, 2H), 7.46 (d, J = 8.1, 2H), 5.88 (bs, 1H), 4.78 (m, 1H), 4.15 (dt, J = 12, 4, 1H), 3.86-3.71 (m, 2H), 3.64 (s, 2H), 3.42-3.34 (m, 2H), 3.04 (s, 3H), 2.85 (m, 2H), 1.62 (s, 9H), 1.61 (s, 9H).
- MS: APCI (+): 609 (M+H)[minor], 497 (-2 tert butyls)[major];
   LC-MS: s, 99 %
   The above 2-(tert-butoxyoxalyl-amino)-7-((2-(4-methanesulfonyl-phenyl)acetylamino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (60 mg, 0.098 mmol) was dissolved in
   50 % trifluoroacetic acid in dichloromethane (2 ml) and allowed to stir at room temperature for 3 hours. The reaction mixture was concentrated in vacuo, the residue titurated with diethyl ether (3x), and dried in vacuo which afforded 45 mg (92 %) of the title compound
- <sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.34 (s, 1H), 8.47 (m, 1H), 7.82 (d, J = 7.8, 2H), 7.50 (d, J = 7.8, 2H), 4.75 (bs, 1H), 4.10 (m, 1H), 3.69 (m, 1H), 3.60 (d, J = 3.6, 2H), 3.52 (m, 1H), 3.35 (m, 2H), 3.18 (s, 3H), 2.83 (m, 2H).

MS: APCI (-): 495 (M-H); LC-MS: s, 95 %.

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as a solid.

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# 2-((3-Carboxy-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl)carbamoyl)nicotinic acid

2-(tert-Butoxyoxalyl-amino)-5-aminomethyl-4,7-dihydro-5H-thieno[2,3-5 c]pyran-3-carboxylic acid tert-butyl ester (164 mg, 0.58 mmol) was stirred for 20 h at 80 °C with furo[3,4-b]pyridine-5,7-dione (86.1 mg, 0.58 mmol) in a mixture of tetrahydrofuran (1.0 ml) and N,N-dimethylformamide (0.25 ml). The volatiles were removed in vacuo and the residue was dissolved in ethyl acetate (50 ml) and washed with water (3 x 30ml). The organic layer 10 was dried(MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue (78 mg) was purified by preparative TLC (hexane/ethyl acetate, 50:50) which afforded 2 products: 2-((2-amino-3-tert-butoxycarbonyl-4,7dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl)carbamoyl)nicotinic acid (A) (27.9 mg, 11 %) and 2-amino-5-(5,7-dioxo-5,7-dihydro-pyrrolo[3,4-15 b]pyridin-6-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (B) (21.3 mg, 9 %).

(A)

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>): δ 9.02 (s, 1H), 8.74 (d, J = 3.3, 1H), 8.14 (d, J = 7.5, 1H), 7.40 (dd, J = 4.8, J = 5.1, 1H), 6.71 (m, 1H), 5.98 (s, 2H), 4.63 (s, 2H), 4.00 (m, 1H), 3.42 (m, 1H), 2.90 (dd, J = 3.3, J = 3.6, 1H), 2.59 (dd, J = 11, J = 11, 1H), 1.48 (s, 9H). MS m/z 434 (M+);

25 **(B)**<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  8.99 (d, J = 5.1, 1H), 8.20 (d, J = 9, 1H), 7.64 (dd, J = 5.7, 4.8, 1H), 5.94 (s, 2H), 4.60 (d, J = 14, 1H), 4.51 (d, J = 14, 1H), 4.05 (m, 2H), 3.87 (d, J = 12.5, 1H), 2.92 (d, J = 17, 1H), 2.61 (m, 1H), 1.53 (s, 9H).

30 MS: APCI (+): 416 (M+1)[minor], 360 (M- tert-butyl) [major].

To a solution of the above 2-((2-amino-3-*tert*-butoxycarbonyl-4,7-dihydro-<sup>3</sup> 5H-thieno[2,3-c]pyran-5-ylmethyl)carbamoyl)nicotinic acid (27.9 mg, 0.064 mmol) in tetrahydrofuran (2 ml) was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (38 mg, 0.193 mmol) and triethylamine (9 μl, 0.064 mmol).

- 5 The resulting mixture was stirred at room temperature for 20 h. The solvent was removed in vacuo and the residue was dissolved in dichloromethane (20 ml) and washed with water (3 x 10 ml). The extracts were dried(MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was purified by preparative TLC (0.5mm, hexane/ethyl acetate,
- 1/1 to 2/3 gradient). After evaporation of the solvent <u>in vacuo</u> 917 mg (46 %) of 2-(3-*tert*-butoxycarbonyl-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl)carbamoyl)nicotinic acid was isolated as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>): δ 9.04 (s, 1H), 8.75 (s, 1H), 8.15 (d, J = 7.5, 1H), 7.42 (dd, J = 6.9, J = 5.1, 1H), 6.73 (m, 1H), 4.81 (dd, J = 15.3, J = 14.4, 2H), 4.03 (m, 1H), 3.83 (m, 1H), 3.47 (m, 1H), 2.99 (d, J = 17.1, 1H), 2.59 (dd, J = 11.1, J = 10.8, 1H), 1.61 (s, 9H), 1.48 (s, 9H).

The above 2-(3-*tert*-butoxycarbonyl-2-(*tert*-butoxyoxalyl-amino)-4,720 dihydro-5H-thieno[2,3-c]pyran-5-ylmethyl)carbamoyl)nicotinic acid (13.1 mg, 0.023 mmol) was stirred in 50 % trifluoroacetic acid in dichloromethane (2 ml) at room temperature for 7 h. The solvent was evaporated <u>in vacuo</u> which afforded 10 mg (96%) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>): δ 9.04 (s, 1H), 8.77 (d, J = 7.7, 1H), 8.16 (d, J = 7.5, 1H), 7.60 (d, J = 7.8, 1H), 4.88 (d, J = 9, 1H), 4.76 (d, J = 9, 1H), 3.96 (m, 1H), 3.02 (m, 1H), 2.78 (m, 1H). MS: 481 (M+33).

MS: 506 (M-55).

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7-(2,4-Dioxo-5-pyridin-2-ylmethylene-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a mixture of 2-(*tert*-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (1.0 g, 2.42 mmol), 5-pyridin-2-ylmethylene-thiazolidine-2,4-dione (0.55 g, 2.66 mmol, prepared in a similar way as described in J. *Med. Chem.* **41** (10), 1619-1630 (1998)) and triphenylphosphine (0.7 g, 2.66 mmol) in dry tetrahydrofuran (75 ml) cooled to 0 °C under a nitrogen atmosphere was added diethyl azodicarboxylate (DEAD) (420 μl ml, 2.66 mmol). The reaction mixture was allowed to stir overnight, slowly warming to room temperature. The volatiles were evaporated in vacuo, the resultant solid was washed with diethyl ether, filtered off and dried in vacuo at 50 °C for h affording 1.4 g (96 %) of 2-(*tert*-butoxyoxalyl-amino)-7-(2,4-dioxo-5-pyridin-2-ylmethylene-thiazolidin-3-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid.

TLC:  $R_f = 0.46$  (ethyl acetate/heptane 1:1).

The above di-*tert*-butyl ester ( 1.0 g, 1.66 mmol) was dissolved in 25 % trifluoroacetic acid in dichloromethane (30 ml). The reaction was stirred at room temperature for 16 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with diethyl ether (50 ml). The precipitate was filtered off, washed with diethyl ether, dried <u>in vacuo</u> at 50 °C for 16 h which afforded 0.8 g of semi pure <u>title compound</u>. The <u>title compound</u> (0.8 g) was suspended in ethyl acetate (25 ml) and heated at reflux temperature for 0.5 h. Isopropanol (5 ml) was added and the mixture was cooled to room temperature the precipitate filtered off and dried <u>in vacuo</u> at 50 °C for 16 h which afforded 0.37 g (37 %) of the <u>title compound</u> as a solid.

Calculated for C<sub>20</sub>H<sub>15</sub>N<sub>3</sub>O<sub>8</sub>S<sub>2</sub>, 0.5 x H<sub>2</sub>O, 0.75 x isopropanol; C, 49.17 %; H, 4.08 %; N, 7.73 %. Found: C, 48.97 %; H, 4.03 %; N, 7.45 %.

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#### **EXAMPLE 37**

# 7-(2,4-Dioxo-5-pyridin-2-ylmethyl-thiazolidin-3-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 5-pyridin-2-ylmethylene-thiazolidine-2,4-dione (5.0 g, 0.024 mol, prepared in a similar way as described in J. Med. Chem. 41 (10), 1619-1630 (1998)) in tetrahydrofuran (300 ml) was added 10 % palladium on carbon (1 g) and the resulting mixture was hydrogenated. After 50 ml of hydrogen was consumed and additional portion of 10 % palladium on carbon (5 g) was added and the hydrogenation was continued at 50 psi for 16 h. The mixture was filtered and the filtrate evaporated in vacuo. The residue was subjected to flash column chromatography (1 l silicagel) using a mixture of ethyl acetate/hexane (1:1) as eluant. Semi pure fractions were collected and the solvent evaporated in vacuo affording 0.8 g (16 %) of 5-pyridin-2-ylmethyl-thiazolidine-2,4-dione as a solid.

To a mixture of 2-(*tert*-butoxyoxalyl-amino)-7-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.7 g, 1.69 mmol), 5-pyridin-2-ylmethyl-thiazolidine-2,4-dione (0.36 g, 1.86 mmol) and triphenylphosphine (0.49 g, 1.86 mmol) in dry tetrahydrofuran (40 ml) cooled to 0 °C under a nitrogen atmosphere was added diethyl azodicarboxylate (DEAD) (290 μl ml, 1.86 mmol). The reaction mixture was allowed to stir overnight, slowly warming to room temperature. The volatiles were evaporated <u>in vacuo</u>, the resultant residue was subjected to

flash column chromatography (0.5 I silicagel) using a mixture of ethyl acetate/hexane (1:2) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 0.6 g (59 %) of 2-(tert-butoxyoxalyl-amino)-7-(2,4-dioxo-5-pyridin-2-ylmethyl-thiazolidin-3-ylmethyl)-4,7-

5 dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as a solid. TLC:  $R_f = 0.43$  (ethyl acetate/heptane 1:1).

The above di-*tert*-butyl ester ( 0.5 g, 0.83 mmol) was dissolved in 25 % trifluoroacetic acid in dichloromethane (25 ml). The reaction was stirred at room temperature for 16 h. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with diethyl ether (20 ml). The precipitate was filtered off, washed with diethyl ether, dried <u>in vacuo</u> at 50 °C for 1 h which afforded 0.3 g of semi pure <u>title compound</u>. The <u>title compound</u> (0.3 g) was suspended in isopropanol (15 ml) and heated at reflux temperature for 5 min., cooled to room temperature and the precipitate filtered off and dried <u>in vacuo</u> at 50 °C for 16 h which afforded 0.2 g (49 %) of the <u>title</u> compound as a solid.

M.p.: > 250 °C;

Calculated for  $C_{20}H_{17}N_3O_8S_2$ , 0.25 x  $H_2O$ ;

20 C, 48.43 %; H, 3.56 %; N, 8.47 %. Found: C, 48.41 %; H, 3.57 %; N, 8.10 %.

#### **EXAMPLE 38**

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7-(5-(4-Methoxy-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl)=2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in Example 37.

M.p.: 236 - 238 °C;

Calculated for  $C_{22}H_{18}N_3O_9S_2$ , 0.5 x  $H_2O$ ;

5 C, 50.09 %; H, 3.63 %; N, 5.31 %. Found:

C, 49.92 %; H, 3.59 %; N, 5.18 %.

# **EXAMPLE 39**

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7-[5-(4-Acetylamino-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl]-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid
The title compound was prepared in a similar way as described in Example 37.

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M.p.: > 250  $^{\circ}$ C;

Calculated for  $C_{23}H_{19}N_3O_9S_2$ , 2 x  $H_2O$ ;

C, 47.50 %; H, 3.99 %; N, 7.23 %. Found:

C, 47.60 %; H, 3.45 %; N, 6.80 %.

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#### EXAMPLE 40

7-[5-(3,5-Dimethoxy-benzylidene)-2,4-dioxo-thiazolidin-3-ylmethyl]-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid
The <u>title compound</u> was prepared in a similar way as described in Example 37.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.37 (s, 1H), 7.92 (s, 1H), 6.80 (d, J = 1.8, 2H), 6.66 (t, J = 2.1, 1H), 5.00 (m, 1H), 4.06 (bm, 2H), 3.81 (s, 6H), 3.71 (dd, J = 6.6, 6, 2H), 2.83 (m, 2H).

MS: APCI (+): 549 (M+H); LC-MS; s, 90 %.

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# EXAMPLE 41

7-[5-(1H-Imidazol-4(5)-ylmethylene)-2,4-dioxo-thiazolidin-3-ylmethyl]-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in Example 37.

20 M.p.: > 250 °C;

Calculated for  $C_{18}H_{14}N_4O_8S_2$ ;

C, 40.65 %; H, 2.56 %; N, 9.17 %. Found:

C, 40.54 %, H, 2.55 %; N, 9.46 %.

## **EXAMPLE 42**

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5-(1,3-Dioxo-4,7-epoxido-1,3,4,5,6,7-hexahydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid To a solution of 2-(tert-butoxyoxalyl-amino)-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (0.20 g, 0.48 10 mmol) in tetrahydrofuran (5 ml) was added 10-oxa-4-azatricyclo(5,2,1,0,2,6)decane-3,5-dione (81 mg, 0.48 mmol) and triphenylphosphine (126 mg, 0.48 mmol). The mixture was cooled to 0  $^{\circ}$ C and diisopropylazodicarboxylate (94.5 µl, 0.48 mmol) was added via syringe. The reaction was stirred for 18h, while slowly warming to room 15 temperature. The volatiles were evaporated in vacuo, and the residue diluted into ethyl acetate (50 ml), washed with saturated sodium bicarbonate (3 x 50 ml), brine (3 x 50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The semi-solid residue was subjected to preparative thin layer chromatography using a mixture of ethyl acetate/hexanes (4:1) as eluant. Fraction with R<sub>f</sub>=0.68 was isolated which 20 afforded 64 mg (24 %) of 2-(tert-butoxyoxalyl-amino)-5-(1,3-dioxo-4,7epoxido-1,3,4,5,6,7-hexahydro-isoindol-2-ylmethyl)-4,7-dihydro-5Hthieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  12.47 (s, 1H), 4.89 (m, 2H), 4.80-4.61 (m, 2H), 3.93-3.86 (m, 1H), 3.83-3.79 (m, 1H), 3.62-3.57 (dd, J = 12.6, 25 3.6, 1H), 2.92 (q, 6.9, 2H), 2.60 (dd, J = 17.1, 10.5, 2H), 1.85 (m, 2H), 1.60 (s, 18H). MS: APCI (-): 561 (M-H).

The above 2-(tert-butoxyoxalyl-amino)-5-(1,3-dioxo-4,7-epoxido-1,3,4,5,6,7-hexahydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-30 c]pyran-3-carboxylic acid tert-butyl ester (51 mg) was dissolved in

50% trifluoroacetic acid in dichloromethane (5ml) and stirred at room temperature for 2 h. The reaction mixture was evaporated <u>in vacuo</u> and the residue titurated with diethyl ether (3 x 10 ml). The solid was filtered of and dried affording 30 mg (71 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 7.68 (bs, 1H), 4.69 (s, 2H), 4.67 (d, J = 15, 1H), 4.56 (d, J = 15, 1H), 3.63 (bm, 1H), 3.50 (d, J = 5, 1H), 3.46 (d, J = 5, 1H), 3.08 (d, J = 15, 2H), 2.94 (d, J = 2.4, 1H), 2.89 (m, 1H), 1.64 (s, 4H).

10 MS: APCI (-): 449 (M-H); LC-MS: s, 95 %

## **EXAMPLE 43**

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7-(((2R)-2-Amino-3-phenyl-propionylamino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid, trifluoroacetic acid salt.

To a stirred solution of a mixture of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester and 2-amino-5-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (4.7 g, 16 mmol) was added diisopropylethylamine (2.8 ml, 16 mmol) and succinimidyl-2,2,2-trichloroethylcarbonate (4.8 g , 16 mmol) portion wise. The reaction mixture was stirred at room temperature for 18 h, washed with saturated sodium hydrogen carbonate, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was chromatographyed-on-silica-(90-g)-using-a mixture-of-ethyl—acetate/heptane (1:1) as eluant. Pure fraction were collected and the solvent evaporated in vacuo affording 6.78 g of crude product which was

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dissolved in dichloromethane (5 ml) followed by heptane (30 ml) which was added as a top layer. After crystallisation and filtration 5.44 g (74 %) of 2-amino-7-((2,2,2-trichloro-ethoxycarbonyl-amino)methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester was obtained as an oil.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 1.55 (s, 9H), 2.78 (m, 2H), 3.32 (m, 1H), 3.62 (m,1H), 3.72 (m,1H), 4.15 (m, 1H), 4,68 (m, 1H), 4.71 (s, 2H), 6.00 (s, 2H). The above 2-amino-7-((2,2,2-trichloro-ethoxycarbonylamino)methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (4.0 g, 8.0 mmol) was dissolved in a mixture of tetrahydrofuran (15 ml) and a aqueous phosphate buffer (pH 3; 5 ml) followed by addition of zinc (16 g, 0.244 mol). The reaction mixture was stirred for 6 h at room temperature at which time the solvent was removed <u>in vacuo</u>. To the residue was added diethyl ether (20 ml) and water (40 ml). Sodium carbonate was added to the aqueous phase until pH = 8 and the aqueous phase was extracted with dichloromethane (3x). The combined organic phases were dried (MgSO<sub>4</sub>), filtered and the solvent removed <u>in vacuo</u>. The residue was purified by flash chromatography on silica gel (90 g) using a mixture of dichloromethane/ethanol/25 % ammonia in water 100:10:0.7 as eluant.

20 Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 1.52 g (61 %) of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.
<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 1.45 (s, 9H), 2.69 (dt, 2H).
Calculated for C<sub>13</sub>H<sub>20</sub>N<sub>3</sub>O<sub>3</sub>S;

25 C, 54.91 %; H, 7.09 %; N, 9.85 %. Found: C, 54.53 %; H, 7.19 %; N, 9.61 %. LC-MS : Mw = 285,2 R<sub>t</sub>= 4.14 min

To a solution of boc-D-phe-OH (0.28 g, 1.05 mmol) in dichloromethane (10 ml) was added 1-hydroxy benzotriazole (0.14 g, 1.05 mmol) and 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (0.18 g, 1.054 mmol). The reaction mixture was stirred for 15 min at room temperature.

2-Amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.30 g, 1.054 mmol) dissolved in dichloromethane (15

ml) was added. Ethyl diisopropylamine (0.18 ml, 1.05 mmol) was added and the reaction mixture was stirred over night at room temperature. The reaction was washed with 10 % aqueous citric acid (15 ml), saturated aqueous sodium hydrogencarbonate, dried (MgSO<sub>4</sub>), filtered and the

solvent removed <u>in vacuo</u> affording 594 mg (100 %) of 2-amino-7-(((1R)-2-*tert*-butoxycarbonylamino-3-phenyl-propionylamino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR(CDCl<sub>3</sub>)  $\delta$  1.42 (s, 9H), 1.55 (s, 9H), 2.73 (m, 2H), 3.05 (m, 2H), 3.16 (m, 1H), 4.06 (m, 1H), 4.32 (m,1H), 5.05 (s, 1H), 6.01 (s, 2H), 6.10 (s, 1H), 7.20 (m, 5H).

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LC-MS: Mw = 532.2,  $R_t = 7.11$ .

2-Amino-7-(((1R)-2-*tert*-butoxycarbonylamino-3-phenyl-propionylamino)-methyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.58 g, 1.09 mmol) was dissolved in dichloromethane (15 ml).

Triethylamine (0.3 ml, 2.18 mmol) was added and the reaction mixture was cooled with in a ice bath. Imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.43 g, 2.18 mmol) dissolved in dichloromethane (5 ml) was added to the reaction mixture. The reaction mixture was stirred overnight at room temperature diluted with dichloromethane (20 ml), washed with 1 N

hydrochloric acid (15 ml), saturated sodium hydrogencarbonate (15 ml), dried (MgSO<sub>4</sub>), filtered and the solvent removed in vacuo. The residue was purified by flash chromatography silica gel (40 g) using a mixture of ethyl acetate/heptane 1:1 as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 512 mg (69 %) of 7-((1R)-(2-tert-

25 butoxycarbonylamino-3-phenyl-propionylamino)methyl)-2-(*tert*-butoxyoxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 1.42 (s, 9H), 1.59 (s, 9H), 1.61 (s, 9H), 2.86 (m, 2H), 3.02 (m, 2H), 3.15 (m, 1H), 3.64 (m, 1H), 3.87 (m, 1H), 4.09 (m, 1H), 4.28 (m, 1H), 4.51 (m, 1H), 4.67 (m, 1H), 5.10 (s, 1H), 6.00 (s, 1H), 7.20 (m, 5H), 12.5 (s, 1H).

7-((1R)-(2-tert-Butoxycarbonylamino-3-phenyl-propionylamino)methyl)-2-(tert-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyrran-3-carboxylic acid tert-butyl ester (0.51 g, 0.76 mmol) was dissolved in dichloromethane (5 ml). Trifluoroacetic acid (5 ml) was added and the reaction mixture was stirred for 2 h at room temperature. The solvent was removed in vacuo ( stripped 3 times with dichloromethane) which afforded 314 mg (92 %) of the title compound.

Calculated for C<sub>20</sub>H<sub>21</sub>N<sub>3</sub>O<sub>7</sub>S; 1 x CF<sub>3</sub>COOH, 1 x H<sub>2</sub>O;

10. C, 45.60 %; H, 4.17 %; N, 7.25 %. Found:

C, 45.78 %; H, 4.20 %; N, 7.05 %.

LC-MS: RT=3.61 / RT=3.77 Mw = 448.2

#### **EXAMPLE 44**

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7-((2-Acetylamino-3-(4-hydroxy-phenyl)-propionylamino)-methyl)-2-(oxalylamino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

To a mixture of Ac-D-Tyr-OH (235 mg, 1.05 mmol) dissolved in dichloromethane (10 ml) was added 1-hydroxybenzotriazole (0.14 g, 1.05 mmol), 1-ethyl-3-(3-dimethylamino propyl)carbodiimide hydrochloride (0.20g, 1.05 mmol) and the reaction mixture was stirred for 15 min at room temperature. 2-Amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid *tert*-butyl ester (0.3 g, 1.05 mmol) dissolved in 25 dichloromethane (10 ml) was added followed by N,N-diisopropylethylamine (0.18 ml, 1.05 mmol). The resulting reaction mixture was stirred for 18-h-at-room temperature, diluted with dichloromethane (15-ml) was washed with 10 % aqueous citric acid (25 ml), saturated sodium hydrogencarbonate, dried (MgSO<sub>4</sub>), filtered and the solvent removed in

vacuo. The residue was purified by flash chromatography on silica gel (40 g) using ethyl acetate as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 304 mg (59 %) of 7-((2-acetylamino-3-(4-hydroxy-phenyl)propionylamino)methyl)-2-amino-4,7-dihydro-5H-

thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) double set of peaks from diastereomers; selected peaks: δ 1.55 (s, 9H), 1.95 (s, 3H), 2.74 (m, 2H), 2.92 (m, 2H), 3.23 (m, 1H), 3.63 (m, 2H), 6.05 (s, 2H).

LC-MS:  $R_t = 5.17$ , Mw = 490.4

7-((2-Acetylamino-3-(4-hydroxy-phenyl)propionylamino)methyl)-2-amino-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.3 g, 0.61 mmol) was dissolved in dichloromethane (15 ml). Triethylamine (0.17 ml, 1.22 mmol) was added and the reaction mixture was cooled to 0° C. Imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.24, 1.22 mmol)

dissolved in dichloromethane (10 ml) was added dropwise. The resulting reaction mixture was stirred at room temperature for 18 h.

Dichloromethane (20 ml) was added and the mixture was washed with 1 N hydrochloric acid (15 ml), saturated sodium hydrogencarbonate (20 ml), dried (MgSO<sub>4</sub>), filtered and the solvent removed in vacuo. The residue was purified by flash chromatography on silica gel (40 g) using ethyl

acetate as eluant. Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 208 mg (55 %) of 7-((2-acetylamino-3-(4-hydroxy-phenyl)-propionylamino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

LC-MS: Mw = 618.4, R<sub>t</sub> = 6.97
 7-((2-Acetylamino-3-(4-hydroxy-phenyl)-propionylamino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.2 g, 0.32 mmol) was dissolved in dichloromethane (8 ml) and trifluoroacetic acid (4 ml) was added. The reaction mixture was stirred
 7 h at room temperature. The solvent was evaporated <u>in vacuo</u> (stripped 3 times with dichloromethane) which afforded 200 mg (100 %) of the <u>title</u> compound.

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Calculated for C<sub>22</sub>H<sub>23</sub>N<sub>3</sub>O<sub>9</sub>S, 3 x H<sub>2</sub>O;
 C, 47.22 %; H, 5.22 %; N, 7.51 %. Found:
 C, 47.05 %; H, 4.88 %; N, 7.39 %.

LC-MS:  $R_t = 3.64$ , Mw = 506.4.

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### **EXAMPLE 45**

10 <u>7-((2-Acetylamino-3-methyl-butyrylamino)methyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid.</u>

To a solution of Ac-D-Val-OH (0.17 g, 1.09 mmol) dissolved in dichloromethane (15 ml) was added N,N-dimethylformamide (1 ml), 1-hydroxybenzotriazole (0.15 g, 1.09 mmol) and 1-ethyl-3-(3-dimethylamino-propyl)carbodiimide hydrochloride (0.21 g, 1.09 mmol). The reaction mixture was stirred for 15 min. at room temperature at which time a solution of 2-amino-7-aminomethyl-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.31 g, 1.09 mmol) in dichloromethane (10 ml) was added followed by N-N-diisopropylethylamine (0.186 ml, 1.09 mmol). The resulting mixture was stirred over night at room temperature diluted with dichloromethane (10 ml) washed with 10 % aqueous citric acid (20 ml), sodium hydrogencarbonate, dried (MgSO<sub>4</sub>), filtered and the solvent was evaporated in vacuo affording 415 mg (90 %) of 7-((2-acetylamino-3-methyl-butyrylamino)methyl)-2-amino-4,7-dihydro-5H-

thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 0.88 (t, 3H), 0.98 (t, 2H), 1.55 (s, 9H), 2.02 (d, 1H), 2.77 m, (2H), 3.40 (m, 1H), 4.14 (m, 1H).

 $LC-MS: R_t = 5.17 Mw = 426.4$ 

To a mixture of 7-((2-acetylamino-3-methyl-butyrylamino)methyl)-2-amino-30 4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.4 g,

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0.94 mmol) dissolved in dichloromethane (10 ml) and triethylamine (0.26 g, 1.87 mmol) cooled to 0° C was added a solution of imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.37 g, 1.87 mmol) in dichloromethane (10 ml). The resulting mixture was stirred for 18 h at room temperature diluted with dichloromethane (20 ml) washed with 1N hydrochloric acid (15 ml), saturated sodium hydrogencarbonate, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo which afforded 515 mg (97 %) of 7-((2-acetylamino-3-methyl-butyrylamino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

LC-MS:  $R_t = 7.11$ , Mw = 554.4.

HPLC:  $R_t = 34.16$ , Area (%) = 100 %.

To a solution of the above 7-((2-acetylamino-3-methyl-butyrylamino)-methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.5 g, 0.90 mmol) dissolved in dichloromethane (3 ml) was added trifluoroacetic acid (1 ml) and the reaction mixture was stirred for 18 h at room temperature. Trifluoroacetic acid (4 ml) was added and the mixture was stirred for an additional 3 hours at room temperature. The volatiles were evaporated <u>in vacuo</u> (and stripped 3 times with dichloromethane) affording 282 mg (71 %) of the <u>title</u> compound.

Calculated for C<sub>18</sub>H<sub>23</sub>N<sub>3</sub>O<sub>8</sub>S, 2 x H<sub>2</sub>O; C, 45.28 %; H, 5.70 %; N, 8.80 %. Found: C, 45.20 %; H, 5.50 %; N, 8.80 %.

25 LC-MS:  $R_t = 3.60$ , Mw = 442.2

# **EXAMPLE 46**

2-(Oxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The title compound was prepared in a similar way as described in Example 25.

M.p.: 210 - 212 °C;
 Calculated for C<sub>18</sub>H<sub>14</sub>N<sub>2</sub>O<sub>9</sub>S<sub>2</sub>, 0.5 x H<sub>2</sub>O, 0.75 x Ethyl acetate;
 C, 44.49 %; H, 3.83 %; N, 5.32 %. Found:
 C, 44.70 %; H, 3.61 %; N, 4.90 %.

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# EXAMPLE 47 O OH S O OH

2-(Oxalyl-amino)-7-(3-oxo-3H-benzo[d]isoxazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The title compound was prepared in a similar way as described in

15 Example 25.

M.p.: 236 - 237 °C;

Calculated for  $C_{18}H_{14}N_2O_8S$ , 0.3 x  $H_2O$ ;

C, 51.02 %; H, 3.47 %; N, 6.61 %. Found:

C, 51.16 %; H, 3.47 %; N, 6.31 %.

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## **EXAMPLE 48**

5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 6-ethyl ester

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To a solution of 1,4-dioxa-8-aza-spiro[4,5]decane (51.5 g, 0.36 moles) in a mixture of dichloromethane (500 ml) and saturated sodium bicarbonate (500 ml) was added di-*teit*-butyldicarbonate (69.8 g, 0.32 moles) and the reaction was vigorously stirred for 3 hours. and the layers separated. The organic layer was washed with 1N hydrochloric acid (2 x 150 ml), brine (100 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u> affording 75.5 g (97 %) of 1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester as a crystallizing oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>): δ 3.96 (s, 4H), 3.49 (bm, 4H), 1.65 (bm, 4H), 1.45 (s, 9H).

To the above 1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester (4.0 g, 16.4 mmol) dissolved in dry diethyl ether (32 ml) was added 2,2' bipyridyl (1 mg) and the solution was cooled to -75 °C. Tetramethylethylenediamine (3.2 ml, 21.4 mmol) was added followed by dropwise addition of sec-butyl lithium (16.4 ml, 21.4 mmol, 1.3M in cyclohexane). The mixture was stirred at -75 °C for 10 min, then slowly warmed to -20 °C and stirred at that temperature for 0.5 h, then cooled to -30 °C. At this temperature, formaldehyde was generated by heating paraformaldehyde at 150 °C and passed through the mixture with dry nitrogen until the color faded to off-white, at which time water (40 ml) was added. The layers were separated, and the aqueous layer was washed diethyl ether (2 x 50 ml). The combined organic extracts were washed 1N hydrochloric acid (2 x 75 ml), saturated sodium bicarbonate solution (50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue (2.9 g) was purified by silica gel chromatography (hexane/ethyl acetate, 10 % ethyl acetate to 30 % ethyl acetate, gradient). Pure fractions were collected and the solvent evaporated in vacuo affording 1.3 g (29 %) of 7-hydroxy-methyl-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid tert-butyl ester as a thick oil. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 4.42 (bm, 1H), 4.08-3.96 (m, 5H), 3.96-3.88 ( m, 1H), 3.78-3.70 (m, 1H), 3.30-3.16 (bm, 1H), 2.30-1.98 (bs, 1H), 1.96-1.78 (m, 2H), 1.74-1.64 (m, 2H), 1.49 (s, 9H).

To 7-hydroxy-methyl-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester (0.4 g, 1.5 mmol) dissolved in dry tetrahydrofuran (20 ml)

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was added phthalimide (0.28 g, 1.9 mmol), triphenylphosphine (0.5 g, 1.9 mmol) and the mixture was cooled to 0  $^{\circ}$ C in an ice bath.

Diethylazodicarboxylate (0.29 ml, 1.82 mmol) was added dropwise and the mixture was stirred at 0 °C for 0.5 h, then at ambient temperature for 18 h. The solvent was removed in vacuo and the residue was purified by silica gel chromatography (hexane/ethyl acetate, 18 % ethyl acetate to 25 % ethyl acetate, gradient). Pure fractions were collected and the solvent evaporated in vacuo affording 0.29 g (48 %) of 7-(1,3-dioxo-1,3-dihydro-

10 tert-butyl ester.

 $^{1}$ H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.94-7.80 (bs, 2H), 7.80-7.64 (bd, 2H), 4.96-4.70 (2bs, 1H), 4.66-4.52 (m, 1H), 4.30-4.14 (bm, 1H), 4.12-4.04 (m, 2H), 4.04-3.94 (m, 2H), 3.56-3.32 (m, 2H), 2.04-1.92 (m, 1H), 1.90-1.60 (m, 4H), 1.22-1.0 (2bs, 9H).

isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid

15 MS: (M + 1) = 403, (M - Boc) = 303.

To the above 7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester (1.1 g, 2.7 mmol) dissolved in dichloromethane (6 ml) was added 1.0 N hydrogen chloride in diethyl ether (50 ml) and the solution kept at ambient temperature for 62

- h. The precipitate was filtered off and washed with diethyl ether and dried with nitrogen which afforded 0.83 g (90 %) of 2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1,3-dione hydrochloride as a solid.

  <sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  9.2-8.8 (2bs, 2H), 7.8-8.1 (m, 2H), 4,1-3.6 (m, 5H), 2.9 (bs, 1H), 2.2-1.6 (m, 5H).
- 25 MS: (M + 1) = 303.5.

To a suspension of the above 2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1,3-dione hydrochloride (0.7 g, 2.1 mmol) and ethyl chloroformate (0.24 ml, 2.5 mmol) in dry tetrahydrofuran (14 ml) cooled in an ice bath under nitrogen was added diisopropylethylamine (0.95 ml, 5.4 mmol) and the reaction mixture was stirred at ambient temperature for 3 hours. The volatiles were removed in vacuo and the residue was partitioned between dichloromethane (25 ml) and 1N hydrochloric acid (25 ml). The layers were separated, and the aqueous layer extracted with

dichloromethane (20 ml). The combined organic extracts were washed with a saturated sodium bicarbonate solution (50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was triturated with n-butylchloride, filtered and dried with nitrogen which afforded 0.47 g

5 (61 %) of 7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid ethyl ester.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.9 (s, 2H), 7.7(s, 2H), 4.9-4.7 (2bs, 1H), 4.7-4.5 (m, 1H), 4.3-3.9 (m, 5H), 3.9-3.6 (bs, 1H), 3.6-3.3 (m, 2H), 2.0-1.9 (m, 1H), 1.9-1.5 (m, 4H), 1.1-0.7 (2bs, 3H).

10 MS: (M-1) = 373.

A solution of the above 7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid ethyl ester (0.44 g, 1.2 mmol) in a mixture of 1N hydrochloric acid (18 ml) and tetrahydrofuran (18 ml) was heated a 75 °C under nitrogen with stirring for 18 h. The

tetrahydrofuran was removed <u>in vacuo</u> and the residue was extracted with dichloromethane (2 x 75 ml). The combined organic extracts were washed with a saturated sodium bicarbonate solution (50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent removed <u>in vacuo</u> affording 0.42 g (> 100 %) of 2-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1-carboxylic

20 acid ethyl ester as a solid.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.9 (s, 2H), 7.8 (s, 2H), 5.3-5.0 (bm, 1H), 4.6-4.2 (bm, 1H), 4.0 (m, 2H), 3.8-3.6 (bm, 3H), 2.8 (m, 1H), 2.7-2.4 (bm, 3H), 1.0 (bs, 3H).

MS: (M+1)= 330.56.

A mixture of the above 2-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1-carboxylic acid ethyl ester (0.39 g, 1.2 mmol), *tert*-butyl cyanoacetate (0.22 g, 1.55 mmol), sulfur (42 mg, 1.3 mmol) in ethanol (1.5 ml) was degassed. To this mixture, under nitrogen, morpholine (205 μl) was added and the mixture was heated a 50 °C for 13 hours. The solvent was removed in vacuo. The residue (0.74 g) was purified by silica gel chromatography-using a mixture of hexane/ethyl acetate (7:3) as eluant. Pure fraction were collected and the solvent evaporated in vacuo. The residue (0.29 g) was titurated with acetonitrile, filtered, and dried with

nitrogen affording 84 mg (15 %) of 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 3-*tert*-butyl ester 6-ethyl ester.

<sup>1</sup>H MNR (400 MHz, CDCl<sub>3</sub>) δ 7.9-7.7 (2m, 4H), 6.0 (bs, 2H), 5.1-4.8 (bm, 1H), 4.8-4.5 (m, 1H), 4.5-4.2 (m, 1H), 4.1-3.4 (3m, 4H), 3.0 (m, 2H), 1.8-1.4 (m, 10H), 1.1-0.9 (m, 3H).

MS: (M + 1) = 486.

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To the above 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 3-*tert*-butyl ester 6-ethyl ester (48 mg, 0.1 mmol) dissolved in dry tetrahydrofuran (1 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.4 ml) and the solution stirred for 18 h. at ambient temperature. The solvent was removed in vacuo and the residue was dissolved in dichloromethane (25 ml) and a saturated sodium bicarbonate solution (25 ml) was added. The layers were separated and the aqueous layer was extracted with dichloromethane (25 ml). The combined organic extracts were dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue (63 mg) was dissolved in ethyl acetate and passed through 1 g of silica gel and the solvent evaporated in vacuo affording 55 mg (90 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 3-*tert*-butyl ester 6-ethyl ester as a solid.

The above 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid 3-*tert*-butyl ester 6-ethyl ester (55 mg, 0.09 mmol) was dissolved in 50 % trifluoroacetic acid in dichloromethane (2 ml) and stirred at ambient temperature for 18 h. The volatiles were removed <u>in vacuo</u> and the residue was purified by preparative hplc (column: Kromasil C18, 250 x 4.6 mm., flow: 2 ml/min., gradient: acetonitrile/water, 20 % acetonitrile to 60 % acetonitrile over 20 min.) affording after evaporation <u>in vacuo</u> 13.8 mg

(31 %) of the <u>title compound</u> as a solid. (Kromasil<sup>™</sup> available from e.g. Richard Scientific Inc, Novato CA.

 $^{1}H$  NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  14-13 (bs, 1H), 12.4 (s, 1H), 7.9 (s, 4H), 4.9 (m, 2H), 4.4 (m, 1H), 4.0-2.8 (m, 13H), 0.8 (m, 3H).

5 MS: (M + 1) = 502.

# **EXAMPLE 49**

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5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester (353 mg, 0.88 mmol) was cooled in an ice bath and then dissolved in a solution of 20 % trifluoroacetic acid/dichloromethane (7 ml). The reaction was stirred for 5 minutes in the ice bath then another 3 hours. at ambient temperature, after which it was concentrated in vacuo affording a solid residue. To the solid was added 2N hydrochloric acid (9 ml) and the mixture was heated at 50 °C (oil bath) with stirring for 24 h. The cooled reaction mixture was quenched with saturated sodium bicarbonate solution until the pH was basic. The aqueous layer was extracted with chloroform (3 x 20 ml) and the combined organic extracts dried (K<sub>2</sub>CO<sub>3</sub>), filtered, and the solvent evaporated in vacuo to give 205 mg (91 %) of 2-(4-oxo-piperidin-2-ylmethyl)-isoindole-1,3-dione as a solid.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.90-7.83 (m, 2H), 7.78-7.71 (m, 2H), 3.81-3.73 (m, 2H), 3.43-3.35 (m, 1H), 3.30-3.22 (m, 1H), 2.83 (dt, J = 13, 3, 1H), 2.46 (d, J = 15, 1H), 2.42-2.32 (m, 2H), 2.21 (dd, J = 14, 13, 1H).

30 APCI-MS:  $[M+H]^{+} = 259$ 

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The above 2-(4-oxo-piperidin-2-ylmethyl)-isoindole-1,3-dione (0.20 g. 0.76 mmol) was dissolved in dichloromethane (5 ml). Saturated sodium bicarbonate solution (5 ml) was added followed by di-tert-butyl dicarbonate (0.20 g, 0.91 mmol). The reaction was stirred vigorously for 16 h. after which the organic phase was separated. The aqueous layer was extracted with dichloromethane (2 x 10 ml) and the combined organic extracts were dried(Na<sub>2</sub>SO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (0 to 10% gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 0.23 g (85) %) of 2-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1carboxylic acid tert-butyl ester.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.86 (bs, 2H), 7.72 (bs, 2H), 5.15-4.98 (m, 1H), 4.23-4.14 (m, 1H), 3.90 (t, J = 12, 1H), 3.61-3.52 (m, 2H), 2.78-2.70 (m, 1H), 2.57-2.39 (m, 3H), 1.15 (s, 9H)  $APCI-MS: [M+H]^{+} = 359$ 

The above 2-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1-carboxylic acid tert-butyl ester (0.43 g, 1.2 mmol) was dissolved in absolute ethanol (9 ml). To the solution was added sulfur (42 mg, 1.32 mmol) and tert-butyl cyanoacetate (0.22 g, 1.56 mmol). The mixture was placed under nitrogen and stirred in a 50 °C oil bath. Morpholine (0.21 ml, 2.4 mmol) was added and the reaction was stirred for 16 h. The precipitate formed was filtered off and washed with acetonitrile (2 x 3 ml) and dried which afforded 0.18 g of 2-amino-5-(1,3-dioxo-1,3-dihydroisoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6dicarboxylic acid di-tert-butyl ester (A) (30 %). The filtrate was concentrated in vacuo and the residue purified by silica gel chromatography using a gradient of ethyl acetate/hexane (1:4 to 1:3 30 ~ gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 0.3 g of a mixture of 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3.6-dicarboxylic acid di-tert-butyl ester and 2-amino-7-(1,3-dioxo-1,3dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester. HPLC purification of a small portion of the mixture gave 28 mg of pure 2-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (B).

(A):

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.87-7.82 (m, 2H) 7.73-7.66 (m, 2H), 6.00 (bs, 2H), 5.02-4.87 (m, 1H), 4.72-4.21 (m, 2H), 4.03-3.93 (m, 1H), 3.51 (t, J = 14, 1H), 2.97-2.91 (m, 2H), 1.56 (s, 9H), 1.12-1.09 (s, 9H).

10. LC-MS:  $R_t$ =3.96 min,  $[M+H]^+$  = 514.4

(B):

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<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.88-7.82 (m, 2H), 7.74-7.66 (m, 2H), 5.39-5.19 (m, 1H), 4.30-4.02 (m, 2H), 3.78-3.70 (m, 1H), 3.33-3.18 (m, 1H), 2.86 (dd, *J* = 18, 4, 1H), 2.75-2.61 (m, 1H), 1.54 (s, 9H), 1.13-1.05 (s, 9H). *LC-MS*: R<sub>t</sub>=4.01 min, [M+H]<sup>†</sup> = 514.4

To a solution of the above 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (50 mg, 0.097 mmol) in dichloromethane (3 ml) was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (60 mg, 0.29 mmol). The reaction was placed under nitrogen and stirred for 3 hours. at ambient temperature. The solution was concentrated in vacuo and the residue purified by silica gel chromatography using a 5 % mixture of ethyl acetate/dichloromethane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 54 mg (87 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester.

1 H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  12.52 (s, 1H), 7.85 (bs, 2H), 7.74-7.67 (m, 2H), 5.08-4.92 (m, 1H), 4.93-4.40 (m, 2H), 3.97-3.87 (m, 1H), 3.53 (t, J = 14, 1H), 3.11-2.99 (m, 2H); 1.62 (s, 18H), 1.14-1.12 (2s, 9H)... *APCI-MS*: [M-H] = 641

The above 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (54 mg, 0.084 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (2 ml). The reaction was stirred at ambient temperature for 7 h., concentrated in vacuo and the residue evaporated in vacuo from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane, filtered off and dried in vacuo which afforded 41 mg (90 %) of the title compound as a solid.

1 H NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  12.31 (s, 1H), 9.36 (bs, 2H), 7.93-7.90 (m, 2H), 7.88-7.85 (m, 2H), 4.43 (d, J = 16, 1H), 4.26 (d, J = 16, 1H), 4.03-3.91 (m, 2H), 3.83-3.76 (m, 1H), 3.31 (dd, J = 18, 4, 1H), 2.82 (dd, J = 18, 10, 1H).

APCI-MS:  $[M+H]^{+} = 430$ 

HPLC (254.4nm): R<sub>t</sub>=6.72 min, 98 %

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#### **EXAMPLE 50**

(L)-5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of L-aspartic acid (120 g, 0.90 mol) in methanol (600 ml) cooled to -20 °C was added thionylchoride (93 ml, 1.29 mol) dropwise over 0.5 h. The cooling bath was removed and the mixture was stirred for 1 h, before diethyl ether (1.8 L, containing 50 ml 1 N hydrochloric acid in diethyl ether) was added upon cooling. The resulting precipitate was filtered off and washed with diethyl ether. The product was recrystallized twice:

First recrystallization: The product was dissolved in warm methanol-(600 ml) and reprecipitated with 1.8 ml diethyl ether (containing 50 ml 1 N hydrochloric acid in diethyl ether).

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Second recrystallization: The product was dissolved in warm methanol (250 ml) and reprecipitated with 1.0 m diethyl ether (containing 50 ml 1 N hydrochloric acid in diethyl ether).

This afforded 75 g ( 45 %) of L-aspartic acid  $\beta$ -methyl ester hydrochloride as a solid.

To a solution of the above  $\beta$ -methyl ester (50 g, 0.27 mol) in water (120 ml) cooled to 0 °C was added triethylamine (95 ml, 0.68 mol) and methyl acrylate (74 ml, 0.82 mol). The reaction mixture was stirred for 3 hours before the cooling bath was removed. After stirring for an additional 1 h the mixture was washed with petrol ether (2 x 400 ml), before tert-butanol (40 ml) and di-tert-butyl dicarbonate (74 g, 0.34 mol) was added and the reaction mixture was stirred for 16 h. The mixture was washed with petrol ether (2 x 400 ml), cooled to 0 °C and the pH was adjusted to 3 with concentrated hydrochloric acid. After extraction with ethyl acetate (3 x 200 ml) the organic phase was washed with brine (200 ml), dried (MgSO<sub>4</sub>), filtered and the volatiles evaporated in vacuo. The residue was subjected to column chromatography on silicagel using a mixture of ethyl acetate/hexane/methanol/acetic acid (25:25:2.5:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo which afforded 60 g (66 %) of 2-(tert-butoxycarbonyl-(2-methoxycarbonyl-ethyl)amino)-succinic acid 4-methyl ester as a solid. To a solution of the above di-ethyl ester (96.9 g, 0.29 mol) in dry degassed tetrahydrofuran (1.0 l) was added sodium methoxide (161 ml, 30% solution in methanol) and the reaction mixture was refluxed under nitrogen for 16 h with mechanical stirring. The reaction mixture was cooled to room temperature, the volatiles remove in vacuo until a wet cage was observed. Water (500 ml) was added and the reaction mixture was refluxed for 16 h. The remaining organic solvents were evaporated in vacuo before the pH was adjusted to 2.5 with concentrated hydrochloric acid. The aqueous phase was extracted with ethyl acetate (3 x 300 ml) and the combined organic phases were washed with brine (100 ml), dried (MgSO<sub>4</sub>) and filtered. tert-Butyl amine (25.36 g, 0.350 mol) was added

dropwise under stirring whereupon a off white precipitate was formed. The precipitate was filtered off and washed with ethyl acetate, dried <u>in vacuo</u> affording 74.4 g (81 %) of 4-oxo-piperidine-1,2-dicarboxylic acid 1-*tert*-butyl ester, *tert*-butyl amine salt as a solid.

Analytically pure compound can be obtained from recrystallisation of the crude product from ethanol-diisopropyl ether by heating the compound in ethanol (ca 100 ml per 10 g compound) and while still hot diisopropyl ether is added (ca 250 ml per 10 g compound). Yield in recrystallisation is approximately 50 %.

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A solution of the above 4-oxo-piperidine-1,2-dicarboxylic acid 1-tert-butyl ester, tert-butyl amine salt (3.0 g, 9.48 mmol), tert-butyl cyanoacetate (2.01g, 14.22 mmol), sulfur (0.456 g, 14.22 mmol) and diisopropylethylamine (1.64 ml, 9.48 mmol) was heated to 50 °C under nitrogen for 12 h. The orange-yellow solution was allowed to cool to room temperature before a small precipitate was filtered off. The filtrate was evaporated in vacuo and the residue was divided between ethyl acetate (50 ml) and saturated ammonium chloride (100 ml). The aqueous phase was extracted with ethyl acetate (3 x 50 m) and the combined organic phases were washed with brine (50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was subjected to column chromatography using a mixture of petrol ether/ethyl acetate/methanol (8:4:1) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 2.22 g (58 %) of 2-amino-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3,5,6-tricarboxylic acid 3,6-di-tert-butyl ester as a solid.

To a solution of the above 3,5,6-tricarboxylic acid 3,6-di-*tert*-butyl ester

(0.63 g, 1.58 mmol) in dimethoxyethane (10 ml) cooled to -20 °C was added N-methylmorpholine (174 ml, 1.58 mmol) followed by isobutylchoroformate (205 ml, 1.58 mmol) and the reaction mixture was stirred for two min. before a precipitate was filtered off. The precipitate was rapidly washed with dimethoxyethane (2 x 2.5 ml), recooled to -20 °C

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and a solution of sodium borohydride (90 mg, 2.37 mmol) in water (1 ml) was added in one lot. (Caution - gas evolution).

The reaction mixture was stirred until gas evolution ceases (app. 3 min.) and the mixture was poured into water (25 ml) and extracted with ethyl acetate (10 ml), washed with brine (5 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo affording 0.40 g (66 %) of 2-amino-5-hydroxymethyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester as a solid.

To a mixture of the above 2-amino-5-hydroxymethyl-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester (2.00 g, 5.20 mmol), phthalimide (0.92 g, 6.24 mmol) and triphenylphosphine (1.64 g, 6.24 mmol) in dry tetrahydrofuran (30 ml) cooled to 0 °C under a nitrogen atmosphere was added diethyl azodicarboxylate (DEAD) (0.98 ml, 6.24 mmol). The reaction mixture was allowed to stir overnight, slowly warming to room temperature. Next day the reaction mixture was again cooled to 0 °C and phthalimide (0.46 g, 3.12 mmol), triphenylphosphine (0.82 g, 3.12 mmol) and diethyl azodicarboxylate (DEAD) (0.49 ml, 3.12 mmol) was added in sequence and the reaction mixture was allowed to stir overnight, slowly warming to room temperature. The volatiles were evaporated in vacuo and the resultant solid dissolved in dichlorormethane (20 ml). The residue was subjected to flash column chromatography using a mixture of ethyl acetate/hexane (1:2) as eluant. Fractions were collected affording after evaporation in vacuo 1.0 g of the desired compound contaminated with phthalimide. Recrystallization from ethanol gave 0.23 g (9 %) of pure 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester as a solid.

To the above di-*tert*-butyl ester (0.20 g, 0.39 mmol) dissolved in dichloromethane (4 ml) was added a mixture of imidazol-1-yl-oxo-acetic acid *tert* butyl ester (0.23 g, 1.17 mmol) in dichloromethane (1 ml) under nitrogen. The reaction mixture was allowed to stir at room temperature overnight. The reaction mixture was added dichlorormethane (5 ml) and washed with 1 % hydrochloric acid (10 ml), dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and

the organic phase evaporated <u>in vacuo</u> affording 0.25 g (100 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester.

The above tri-tert-butyl ester (0.25 g, 0.39 mmol) was dissolved in 20 % trifluoroacetic acid in dichloromethane (5 ml). The reaction was stirred at room temperature for 24 h. before diethyl ether (5 ml) was added. The precipitate was filtered off, washed with diethyl ether, dried in vacuo to give 0.150 g of a solid. NMR revealed the presence of a trace amount of material arising from incomplete deprotection. 0.100 g of the crude product was redissolved in 20 % trifluoroacetic acid in dichloromethane (5 ml), and stirred at room temperature for 24 h. before diethyl ether (5 ml) was added. The product was filtered off and washed with diethyl ether and dried in vacuo to give 0.05 g (40 %) of the title compound as a solid.

M.p.: dec.> 240° C
Calculated for C<sub>19</sub>H<sub>15</sub>N<sub>3</sub>O<sub>7</sub>S 1/3 C<sub>2</sub>HF<sub>3</sub>O<sub>2</sub> 1/2 H<sub>2</sub>O;
C, 49.58 %; H, 3.46 %; N, 8.82 %. Found:
C, 49.84 %; H, 3.83 %; N, 8.99 %.

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### **EXAMPLE 51**

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7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of pure 2-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (28 mg, 0.057 mmol) in dichloromethane (2 ml) was added midazol-1-yl-oxo-acetic acid *tert*-butyl ester (35 mg, 0.17 mmol). The reaction was placed under nitrogen and stirred for 12 h. at ambient temperature. The volatiles were evaporated in <u>vacuo</u> and the residue was purified by silica gel chromatography using a mixture of ethyl acetate/hexane (1:3) as eluant. Pure fractions were collected and the solvent evaporated in <u>vacuo</u> affording 25 mg (67 %) of 2-(*tert*-butoxyoxalyl-amino)-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester as an oil.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 12.59-12.53 (bs, 1H), 7.89-7.84 (m, 2H), 7.75-7.67 (m, 2H), 5.61-5.41 (m, 1H), 4.36-4.15 (m, 1H), 4.12-4.06 (m, 1H), 3.90-3.82 (m, 1H), 3.34-3.21 (m, 1H), 2.99-2.93 (m, 1H), 2.84-2.68 (m, 1H), 1.62-1.59 (s, 18H), 1.12-1.06 (s, 9H).

The above 2-(*tert*-butoxyoxalyl-amino)-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (25 mg, 0.039 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (1.5 ml). The reaction was stirred at ambient temperature for 7 h., concentrated <u>in vacuo</u> and the residue evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane, filtered off and dried <u>in vacuo</u> to give 41 mg (85 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.32 (s, 1H), 9.48 (bs, 2H), 7.95-7.91 (m, 2H), 7.89-7.84 (m, 2H), 4.89 (s, 1H), 4.15-4.07 (m, 2H), 3.43-3.28 (2m, 2H, partially obscured by water), 3.04 (bs, 2H). LC-MS: R<sub>t</sub>=1.51 min, [M-H] = 428.4

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### **EXAMPLE 52**

5-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-1,4-dioxa-8-aza-spiro[4.5]decane-8-carboxylic acid *tert*-butyl ester (1.55 g, 3.85 mmol) was cooled in an ice bath and then dissolved in a solution of 20 % trifluoroacetic acid/dichloromethane (15 ml). The reaction was stirred and allowed to slowly warm to ambient temperature during 3 hours. The solution was concentrated <u>in vacuo</u> to give crude 2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1,3-dione which was used directly in the following step (assumed 100 % yield).

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 9.26 (bs, 1H), 8.19 (bs, 1H), 7.78-7.75 (m, 2H), 7.74-7.71 (m, 2H), 4.11-3.98 (m, 5H), 3.90-3.79 (m, 3H), 3.26-3.17 (m, 1H), 2.10-2.00 (m, 3H), 1.92-1.88 (m, 1H).

To a suspension of the above 2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1,3-dione (3.85 mmol) in absolute ethanol (25 ml) was added hydrazine (0.36 ml, 11.55 mmol). The reaction was stirred at 80 °C (oil bath) for 6 h., then cooled to ambient temperature and stirred for an additional 12 h. The thick precipitate was filtered off and washed with ethanol. The filtrate was concentrated in vacuo and reconstituted in dichloromethane (20 ml), forming a small amount of a second precipitate which was filtered off. The filtrate was evaporated in vacuo and the resulting oil was dissolved in water (10 ml) and basified with 1N sodium hydroxide until pH = 10. The aqueous layer was extracted with 20 % isopropyl alcohol/chloroform (12 x 40 ml). The combined organic extracts

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were dried ( $K_2CO_3$ ), filtered and the solvent evaporated <u>in vacuo</u> affording 0.42 g (63 %) of (1,4-dioxa-8-aza-spiro[4.5]dec-7-yl)methylamine as an oil. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  3.94 (bs, 4H), 3.11-3.05 (m, 1H), 2.81 (dt, J = 12, 3, 1H), 2.76-2.65 (m, 2H), 2.58-2.50 (m, 1H), 1.70-1.57 (m, 3H), 1.31 (t, J = 12, 1H).

APCI-MS:  $[M+H]^{+} = 173.2$ 

To a solution of 4-hydroxy-isobenzofuran-1,3-dione (0.51 g, 3.09 mmol) in anhydrous N,N-dimethylformamide (7 ml) under nitrogen was added sodium hydride (130 mg, 3.25 mmol). Immediate evolution of gas and bright yellow color was observed. The mixture was stirred for 5 minutes after which benzyl bromide (1.8 ml, 15.45 mmol) was added. The reaction was stirred for 72 h. Saturated sodium bicarbonate (2 ml) was added and the mixture stirred for 2 minutes, diluted in ethyl acetate (35 ml) and washed with saturated sodium bicarbonate (5 ml), 1N hydrochloric acid (5 ml), and brine (2 x 5 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. To the crude material was added hexane and the formed precipitate was filtered off, washed further with hexane and dried <u>in vacuo</u> to give 0.54 g (69 %) of 4-(benzyloxy)-isobenzofuran-1,3-dione as a solid.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  7.74 (t, 1H, J = 8 Hz), 7.54 (d, 1H, J = 8 Hz), 7.47-7.29 (m, 6H), 5.36 (s, 2H).

A solution of (1,4-dioxa-8-aza-spiro[4.5]dec-7-yl)methylamine (0.19 g, 1.1 mmol) and 4-(benzyloxy)-isobenzofuran-1,3-dione (0.27 g, 1.05 mmol) was prepared in a mixture of distilled dichloromethane (3 ml) and anhydrous N,N-dimethylformamide (2.5 ml) under nitrogen. 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.23 g, 1.21 mmol) was added followed by triethylamine (0.46 ml, 3.3 mmol) and the reaction stirred at ambient temperature for 18 h. The solution was concentrated in vacuo and the residue diluted with ethyl acetate (25 ml) and washed with water (5 ml), saturated sodium bicarbonate (5 ml), and brine (5 ml). The organic layer was evaporated in vacuo and the residue

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purified by silica gel chromatography using a mixture of 5 % methanol/dichloromethane/1 % triethylamine as eluant .Pure fractions were collected and the solvent evaporated <u>in vacuo</u> affording 0.22 g (50 %) of 4-benzyloxy-2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1.3-dione as a semi-solid.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.57 (t, J = 8, 1H), 7.48 (d, J = 7, 2H), 7.42-7.29 (m, 4H), 7.18 (d, J = 8, 1H), 5.31 (s, 2H), 3.94-3.90 (m, 4H), 3.65 (d, J = 6, 2H), 3.16-3.09 (m, 1H), 3.07-3.02 (m, 1H), 2.76 (dt, J = 13, 3, 1H), 1.78 (d, J = 12, 1H), 1.64-1.54 (m, 3H), 1.37 (t, J = 12, 1H), 1.08 (t, J = 7, 10 1H).

LC-MS:  $R_t$ =2.59 min,  $[M+H]^+$  = 409.2

To a solution of the above 4-benzyloxy-2-(1,4-dioxa-8-aza-spiro[4.5]dec-7-ylmethyl)-isoindole-1,3-dione (0.22 g, 0.54 mmol) in 1,4-dioxane (4 ml) was added 4N hydrochloric acid (4 ml) and the reaction stirred in a 65  $^{\circ}$ C (oil bath) for 6 h. The mixture was basified with saturated sodium bicarbonate until pH = 8 and extracted with dichloromethane (3 x 20 ml). The combined organic extracts were dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo affording crude 4-benzyloxy-2-(4-oxopiperidin-2-ylmethyl)-isoindole-1,3-dione as an oil. Which was used without further purification or characterization.

The above crude 4-benzyloxy-2-(4-oxo-piperidin-2-ylmethyl)-isoindole-1,3-dione (0.17 g, 0.47 mmol) was dissolved in dichloromethane (4 ml). Saturated sodium bicarbonate (4 ml) was added followed by di-*tert*-butyl dicarbonate (0.11 g, 0.52 mmol). The reaction was stirred vigorously for 16 h., then the layers were separated. The aqueous layer was extracted with dichloromethane (2 x 10 ml) and the combined organic phases were dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was purified by silica gel chromatography using a mixture of ethyl acetate/hexane (1:2) as eluant. Pure fractions were collected and the solvent was evaporated <u>in vacuo</u> affording 0.14 g (64 %) of 2-(4-

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benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  7.57 (bs, 1H), 7.47-7.31 (m, 6H), 7.18 (bs, 1H), 5.34 (s, 2H), 5.03 (bs, 1H), 4.45-4.14 (m, 1H), 3.89 (t, J = 12, 1H), 3.55 (bs, 2H), 2.76-2.71 (m, 1H), 2.57-2.38 (m, 3H), 1.17 (s, 9H).

A solution of 2-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4-oxo-piperidine-1-carboxylic acid *tert*-butyl ester (0.14 g, 0.30 mmol), sulfur (10.6 mg, 0.33 mmol), and *tert*-butyl cyanoacetate (55 mg, 0.39 mmol) in absolute ethanol (4 ml) was stirred at 50 °C (oil bath). Morpholine (53 μl, 0.6 mmol) was added and the reaction placed under nitrogen and stirred for 16 h. The solution was cooled to ambient temperature, concentrated <u>in vacuo</u> and the residue purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (0 to 5 % gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording a mixture of regioisomers 0.15 g (80 %) of 2-amino-5-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester and 2-amino-7-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester which were not separable by chromatography.

To a solution of the above mixture of 2-amino-5-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester and 2-amino-7-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (0.15 g, 0.24 mmol) in distilled dichloromethane (4 ml) under nitrogen was added imidazol-1-yloxo-acetic acid *tert*-butyl ester (0.14 g, 0.72 mmol) and the reaction mixture was stirred at ambient temperature for 1.5 h. The volatiles were evaporated in vacuo and the crude residue was purified by silica gel chromatography using dichloromethane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affording 50 mg of 2-(*tert*-collected and the solvent evaporated in vacuo affor

butoxyoxalyl-amino)-5-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (**A**) and 50 mg of 2-(*tert*-butoxyoxalyl-amino)-7-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (**B**). Another 50 mg remained as a mixture of the two isomers (**A**) and (**B**).

(A):

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.52 (s, 1H), 7.60-7.31 (m, 7H), 7.20-7.10 (m, 1H), 5.33 (s, 2H), 5.05-4.38 (m, 3H), 3.96-3.83 (m, 1H), 3.52-3.41 (m, 1H), 3.01 (bs, 2H), 1.60 (s, 9H), 1.59 (s, 9H), 1.17-1.14 (s, 9H). LC-MS:  $R_t$ =4.93 min,  $[M+H]^+$  = 748.1

(B):

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.58-12.52 (s, 1H), 7.60-7.30 (m, 7H), 7.20-7.10 (m, 1H), 5.60-5.39 (m, 1H), 5.34 (s, 2H), 4.36-4.02 (m, 2H), 3.86-3.75 (m, 1H), 3.33-3.18 (m, 1H), 2.97-2.90 (m, 1H), 2.83-2.68 (m, 1H), 1.60 (s, 9H), 1.58-1.57 (s, 9H), 1.15-1.09 (s, 9H) LC-MS: R<sub>t</sub>=4.93 min, [M+H]<sup>+</sup> = 748.1

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The above 2-(*tert*-butoxyoxalyl-amino)-5-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (50 mg, 0.067 mmol) was dissolved in a mixture of ethyl acetate/ethanol (3 ml, 1:1). Palladium on activated carbon (10 %, 10 mg) was added and the solution degassed and stirred under hydrogen (1 atm.) for 72 h. TLC analysis indicated that the reaction was incomplete. The mixture was filtered through celite and the filter cake washed with hot ethyl acetate. The filtrate was concentrated <u>in vacuo</u> and purified by silica gel chromatography using a gradient of ethyl acetate/dichloromethane (0 to 5 % gradient) as eluant. Pure fractions were collected and-the solvent-evaporated <u>in-vacuo</u> affording 15 mg (30 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(4-hydroxy-1,3-dioxo-1,3-dihydro-

isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 12.50 (s, 1H), 7.61-7.51 (m, 1H), 7.39-7.34 (m, 1H), 7.17-7.09 (m, 1H), 5.04-4.64 (m, 2H), 4.49-4.34 (m, 1H), 3.90-3.78 (m, 1H), 3.51-3.42 (m, 1H), 3.02 (bs, 2H), 1.60 (s, 18H), 1.17-1.14 (2s, 9H).

The above 2-(*tert*-butoxyoxalyl-amino)-5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-

- dicarboxylic acid di-*tert*-butyl ester (15 mg, 0.023 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (2 ml). The reaction was stirred at ambient temperature for 12 h., concentrated <u>in vacuo</u> and evaporated <u>in vacuo</u> from dichloromethane (3 x 10 ml). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> affording
   6 mg (47 %) of the <u>title compound</u>.
  - <sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.32 (s, 1H), 11.17 (s, 1H), 9.25 (bs, 2H), 7.64 (t, J = 8, 1H), 7.32 (d, J = 8, 1H), 7.24 (d, J = 8, 1H), 4.41-4.23 (m, 2H), 3.96-3.71 (m, 3H), 3.5-3.2 (obscured by water, 1H), 2.83-2.75 (m, 1H).
- 20 *LC-MS*:  $R_t$ =1.53 min,  $[M+H]^{+}$  = 446.2

## **EXAMPLE 53**

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2-(Oxalyl-amino)-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

2-Methyl-benzoic acid methyl ester (1.50 g 10 mmol), N-bromosuccinimide (1.96 g, 11 mmol) and 2,2'-azobis(2-methylpropionitrile) (AIBN) (25 mg,

0.15 mmol) were dissolved in chloroform (3 ml). The solution was heated at reflux for 16 h. cooled and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/hexane (1-2 %) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 2.05 g (89 %) of 2-bromomethyl-benzoic acid methyl ester as a solid.

<sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  7.97 (d, 1H, J = 7.6 Hz), 7.45-7.52 (m, 2H), 7.38 (dt, 1H, J = 1.2, 7.6 Hz), 4.96 (s, 2H), 3.95 (s, 1H).

To a solution of 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (100 mg, 0.20 mmol) and pyridine (0.18 ml, 2.0 mmol) in acetonitrile (1 ml) at room temperature was added benzyl chloroformate (0.28 ml, 2.0 mmol) in 10 aliquots over 48 h. The solution was then taken into ethyl acetate (30 ml), washed with 0.5 N hydrochloric acid (3 x 10 ml), saturated sodium bicarbonate (3 x 10 ml), brine (10 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was evaporated in vacuo. The resulting oil crystallized upon standing for 2 days. The precipitate was filtered off and washed with diethyl ether (3 x 1 ml) affording after drying in vacuo 59 mg (47 %) of 2-benzyloxy-carbonylamino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester as a solid.

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To a solution of 1 N hydrochloric acid in ethyl acetate (1.0 ml) was added 2-benzyloxy-carbonylamino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (52 mg, 0.08 mmol). The solution was stirred at room temperature for 48 h. A precipitate was filtered off which afforded 42 mg (90 %) of 2-benzyloxy-carbonylamino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester hydrochloride as a solid.

<sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  10.60 (s, 1H), 7.60-7.92 (m, 4H), 7.38 (m, 5H), 5.26 (s,

2H), 4.30-5.10 (m, 3H), 3.40-4.00 (m, 2H), 1.57 (m, 9H), 1.15 (m, 9H).

<sup>1</sup>H NMR (DMSO-d<sub>6</sub>): δ 10.45 (s, 1H), 9.40 (s, 1H), 9.25 (s, 1H), 7.89 (m, 4H), 7.39 (m, 5H), 5.22 (s, 2H), 4.39 (d, 1H, J = 15 Hz), 4.28 (m, 1H), 3.95 (m, 2H), 3.79 (m, 1H), 3.20 (m, 1H), 2.70 (m, 1H), 1.48 (s, 9H).

To a solution of the above 2-benzyloxy-carbonylamino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester hydrochloride (42 mg, 0.072 mmol) in ethanol (0.5 ml) was added hydrazine (68 μl, 0.22 mmol). The solution was stirred at 80 °C for 5 h. and at room temperature for 16 h. The mixture was filtered and the filtrate evaporated in vacuo. The residue was extracted with dichloromethane (5 x 1 ml). The combined dichloromethane washes were evaporated in vacuo affording 20 mg (67 %) of 5-aminomethyl-2-benzyloxy-carbonylamino-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

15 <sup>1</sup>H NMR (CDCl<sub>3</sub>): δ 10.55 (bs, 1H), 7.37 (m, 5H), 5.23 (s, 2H), 3.92 (s, 2H),
 2.60-3.10 (m, 3H), 1.53 (s, 9H).

To a solution of the above 5-aminomethyl-2-benzyloxy-carbonylamino-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (20 mg, 0.048 mmol) in acetonitrile (1 ml) at 0 °C was added 20 diisopropylethylamine (18 µl, 0.15 mmol) and 2-bromomethyl-benzoic acid methyl (12 mg, 0.048 mmol). The solution was stirred at 0 °C for 3 hours. and at room temperature for 16 h. Di-tert-butyl dicarbonate (21 mg, 0.096) mmol) was then added to the solution. The solution was then stirred at room temperature for 16 h. The solution was taken into ethyl acetate (30 25 ml), washed with 0.5 N hydrochloric acid (3 x 10 ml), saturated sodium bicarbonate (3 x 10 ml) and brine (10 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was evaporated in vacuo. The solid residue was purified by silica gel chromatography using a 5 % mixture of ethyl acetate/hexane as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording-10-mg-(33-%) of-2-benzyloxycarbonylamino-5-(1-oxo-1,3dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6dicarboxylic acid di-tert-butyl ester as a solid.

<sup>1</sup>H NMR (CDCl<sub>3</sub>): δ 10.59 (s, 1H), 7.81 (m, 1H), 7.52 (m, 1H), 7.39 (m, 7H), 5.25 (s, 1H), 4.22-5.00 (m, 4H), 4.40-4.80 (m, 2H), 2.80-3.10 (m, 2H), 1.55 (s, 9H), 1.25 (s, 9H).

To a solution of the above 2-benzyloxycarbonylamino-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (9 mg, 0.014 mmol) in methanol (2 ml) was added 10 % Pd/C (4 mg). The mixture was stirred under hydrogen (1 atm.) for 3 hours. and then filtered. The filtrate was evaporated in vacuo affording 6 mg (93 %) of 2-amino-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester as a solid.

1 NMR (CDCl<sub>3</sub>): δ 7.80 (m, 1H), 7.50 (m, 1H), 7.44 (m, 2H), 4.22-5.00 (m,

'H NMR (CDCl<sub>3</sub>): δ 7.80 (m, 1H), 7.50 (m, 1H), 7.44 (m, 2H), 4.22-5.00 (m 4H), 4.40-4.80 (m, 2H), 2.80-3.10 (m, 2H), 1.63 (s, 9H), 1.25 (s, 9H).

To a solution of the above 2-amino-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid ditert-butyl ester (6 mg, 0.013 mmol) in acetonitrile (0.5 ml) at room temperature was added imidazol-1-yl-oxo-acetic acid tert-butyl ester (27 mg, 0.13 mmol). The solution was stirred for 3 hours. at room temperature and then diluted with ethyl acetate (20 ml), washed with 0.5 N hydrochloric acid (2 x 5 ml), saturated sodium bicarbonate (2 x 5 ml), brine (5 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of ethyl acetate/hexane (10-25 % gradient) as eluant. Pure fractions were collected and the solvent evaporated in vacuo affording 4 mg (50 %) of 2-(tert-butoxyoxalyl-amino-5-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester as a solid.

<sup>1</sup>H NMR (CDCl<sub>3</sub>): δ 12.49 (s, 1H), 7.80 (m, 1H), 7.50 (m, 1H), 7.44 (m, ---2H), 4.22-5.00-(m, 4H), 4.20-4.90 (m, 2H), 2.90-3.20 (m, 2H), 1.63 (s, 9H), 1.60 (s, 9H), 1.25 (s, 9H).

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### **EXAMPLE 55**

2-(Oxalyl-amino)-7-(1,1,3-trioxo-5-phenyl-1,3-dihydro-isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

- The <u>title compound</u> was prepared in a similar way as described in Example 23 using 2-amino-7-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxo-5-phenyl-1,2-dihydro-1H-isothiazol-3-one as starting material. *O* and *N*-alkylated products were separated by column chromatography.
- <sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 2.85 (bs, 2H), 3.75 (m, 1H), 3.92 (dd, 1H), 4.10 (m, 2H), 5.08 (m, 1H), 7.64 (m, 3H), 7.69 (s, 1H), 7.92 (m, 2H), 12.35 (s, 1H, N*H*CO).

LC-MS:  $R_t = 4.90 \text{ min, m/z: } 493 [M+H]^{+}$ 

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### **EXAMPLE 56**

7-(1,1-Dioxo-5-phenyl-1*H*-isothiazol-3-yloxymethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

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The <u>title compound</u> was prepared in a similar way as described in Example 23-using 2-amino 7-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxo-5-phenyl-1,2-

dihydro-1H-isothiazol-3-one as starting material. O- and N-alkylated products were separated by column chromatography.

 $^{1}$ H-NMR (DMSO-d<sub>6</sub>)  $\delta$  2.86 (bs, 2H), 3.79 (m, 1H), 4.13 (m, 1H), 4.75 (m, 2H), 5.17 (bs, 1H), 7.60 (m, 3H), 7.70 (s, 1H), 7.88 (m, 2H), 12.35 (s, 1H, N*H*CO).

LC-MS:  $\dot{R}_t = 4.78 \text{ min, m/z: } 493 \text{ [M+H]}^+$ 

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# **EXAMPLE 57**

2-(Oxalyl-amino)-5-(1,1,3-trioxo-5-phenyl-1,3-dihydro-isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

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The <u>title compound</u> was prepared in a similar way as described in Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxo-5-phenyl-1,2-dihydro-1*H*-isothiazol-3-one as starting material. *O*- and *N*-alkylated products were separated by column chromatography.

1H-NMR (DMSO-d<sub>6</sub>) δ 2.62 (dd, 1H), 3.05 (d, 1H), 3.88 (m, 2H), 3.98 (m, 1H), 4.60–4.86 (dd, 2H), 7.66 (m, 4H), 7.93 (m, 2H), 12.3 (s, 1H, N*H*CO).

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### **EXAMPLE 58**

5-(1,1-Dioxo-5-phenyl-1*H*-isothiazol-3-yloxymethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-

c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxo-5-phenyl-1,2-dihydro-1*H*-isothiazol-3-one as starting material. *O*- and *N*-alkylated products were separated by column chromatography.

Mp.: 230 - 232 °C;

Calculated for C<sub>20</sub>H<sub>16</sub>N<sub>2</sub>O<sub>9</sub>S<sub>2</sub>, 1xH<sub>2</sub>O;
C, 47.06 %; H, 3.55 %; N, 5.49 %. Found:
C, 46.88 %; H, 3.44 %; N, 5.45 %.

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#### **EXAMPLE 59**

5-(6-Chloro-1,1,3-trioxo-2,3-dihydro-4*H*-thieno[3,2-e]-1,2,4-thiadiazin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

- The <u>title compound</u> was prepared in a similar way as described in Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxide-6-chloro-2,3-dihydro-4*H*-thieno[3,2-e]-1,2,4-thiadiazine-3-one as starting material. *O*-and *N*-alkylated products were separated by column chromatography.
- <sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 2.60 (dd, 1H), 2.98 (d, 1H), 3.87–3.96 (m, 2H), 4.04 (m, 1H), 4.56–4.82 (dd, 2H), 7.0 (s, 1H), 11.95 (s, 1H, N*H*CO), 12.3 (s, 1H, N*H*CO).

5-(6-Chloro-1,1-dioxo-4*H*-thieno[3,2-e]-1,2,4-thiadiazine-3-yloxymethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

The title compound was prepared in a similar way as described in

- Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1,1-dioxide-6-chloro-2,3-dihydro-4*H*-thieno[3,2-e]-1,2,4-thiadiazine-3-one as starting material. *O*-and *N*-alkylated products were separated by column chromatography. Mp.: > 250 °C;
- Calculated for C<sub>16</sub>H<sub>12</sub>ClN<sub>3</sub>O<sub>9</sub>S<sub>3</sub>, 0.75xH<sub>2</sub>O;
   C, 35.89 %; H, 2.54 %; N, 7.85 %. Found:
   C, 35.84 %; H, 2.36 %; N, 7.74 %.

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# EXAMPLE 61

7-(1,3-Dioxo-1,3-dihydro-benzo[d]isothiazol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

- The <u>title compound</u> was prepared in a similar way as described in Example 23 using 2-amino-7-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 1-oxo-1,2-dihydro-1H-benzo[d]isothiazol-3-one as starting material. *O* and *N*-alkylated products were separated by column chromatography.
- 25 LC-MS:  $R_t = 3.82 \text{ min, m/z: } 451 \text{ [M+H]}^{+}$



### **EXAMPLE 62**

5-(1,3-Dioxo-1,3-dihydro-benzo[d]isothiazol-2-ylmethyl)-2-(oxalyl-amino)-

4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid 5

The title compound was prepared in a similar way as described in Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester and 1-oxo-1,2-dihydro-1H-10 benzo[d]isothiazol-3-one as starting material. O- and N-alkylated products were separated by column chromatography.

Mp.: 230 - 231 °C;

Calculated for  $C_{18}H_{14}N_2O_8S_2$ ,  $0.5xH_2O$ ;

C, 47.06 %; H, 3.29 %; N, 6.10 %. Found: 15

C, 46.94 %; H, 3.42 %; N, 6.26 %.

# **EXAMPLE 63**

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5-(5-Benzyl-1,1-dioxo-[1,2,5]thiadiazolidin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid

The title compound was prepared in a similar way as described in 25 Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester and 2-benzyl-

I1\_2\_5]thiadiazolidine 1,1-dioxide as starting material.

LC-MS:  $R_t = 5.00 \text{ min, m/z: } 496 \text{ [M+H]}^{+}$ 

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### **EXAMPLE 64**

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5-(5-Ethyl-1,1-dioxo-[1,2,5]thiadiazolidin-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in

Example 23 using 2-amino-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 2-ethyl-[1,2,5]thiadiazolidine 1,1-dioxide as starting material.

LC-MS:  $R_t = 4.18 \text{ min, m/z: } 434 \text{ [M+H]}^{+}$ 

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### **EXAMPLE 65**

2-(Oxalyl-amino)-7-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

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To a solution of 2-amino-7-aminomethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (100 mg, 0.38 mmol) and *N,N*-

oil.

diisopropylethylamine (72  $\mu$ L, 0.41 mmol) in acetonitrile (6 ml) at 0 °C was added 2-bromomethyl-benzoic acid methyl ester (43 mg, 0.19 mmol). The reaction mixture was stirred for 16 hours and the solvent evaporated in vacuo. The residue was diluted in ethyl acetate (50 ml), washed with 1N hydrochloric acid, saturated sodium bicarbonate, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo, which afforded 50 mg (68 %) of 2-amino-7-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 8 7.86 (d, 1H, J = 8 Hz), 7.55 (t, 1H, J = 8 Hz), 7.45 (t, 2H, J = 8 Hz), 4.88 (dt, 1H. J = 6, 2 Hz), 4.68 (d, 1H, J = 17 Hz), 4.48 (d, 1H, J = 17 Hz), 4.25-4.10 (m, 1H), 4.03 (dd, 1H, J = 17 and J = 3 Hz), 3.80-3.75 (m, 2H), 2.92-2.70 (m, 2H), 1.54 (s, 9H).

To a solution of 2-amino-7-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (50 mg, 0.13 mmol) in tetrahydrofuran (1 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (100 mg, 0.51 mmol). The mixture was stirred at room temperature for 24 hours. The solvent was removed <u>in vacuo</u>. The residue was taken into ethyl acetate (50 ml), washed with saturated sodium
bicarbonate and brine, dried (Na<sub>2</sub>SO<sub>4</sub>) and filtered. The solvent was removed <u>in vacuo</u> and the residue was chromatographed using a mixture of 10% ethyl acetate/dichloromethane as eluent, which afforded 55 mg (83 %) of 2-(*tert*-butoxyoxalyl-amino)-7-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 12.59 (s, 1H), 7.88 (d, 1H, J = 7 Hz), 7.54 (t, 1H, J = 7 Hz), 7.46 (t, 2H, J = 7 Hz), 5.04 (dd, 1H, J = 6 Hz and J = 2 Hz), 4.69 (d, 1H, J = 17 Hz), 4.46 (d, 1H, J = 17 Hz), 4.26-4.10 (m, 2H), 3.77 (dd, 1H, J = 9 Hz and J = 3 Hz), 3.70 (dd, 1H, J = 15 Hz and J = 9 Hz), 3.02-2.80 (m, 30 2H), 1.55 (s, 18H).

4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an

A solution of 2-(*tert*-butoxyoxalyl-amino)-7-(1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl

ester (55 mg, 0.11 mmol) in 50 % trifluoroacetic acid/dichloromethane (2 ml) was stirred for 16 hours. The volatiles were removed in vacuo and the residue was washed with dichloromethane and dried, which afforded 29 mg (50 %) of the title compound as a solid trifluoroacetate.

5 <sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.35 (s, 1H), 7.70 (d, 1H, J = 8 Hz), 7.61 (d, 1H, J = 83 Hz), 7.52-7.47 (m, 2H), 5.04 (s, 1H), 4.59 (d, 1H, J = 18 Hz), 4.58 (d, 1H, J = 18 Hz), 4.19-4.08 (m, 1H), 3.88 (d, 1H, J = 6 Hz), 3.78-3.66 (m, 1H), 3.38 (q, 1H, J = 7 Hz), 2.85 (s, 2H);

LC-MS:  $R_t = 2.12 \text{ min, m/z: } 417 \text{ [M+H]}^+$ 

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# **EXAMPLE 66**

2-(Oxalyl-amino)-5-(2,2,2-trifluoro-acetoxymethyl)-4,7-dihydro-5H-

thieno[2,3-c]pyran-3-carboxylic acid 15

> 2-(tert-Butoxyoxalyl-amino)-5-hydroxymethyl-4,7-dihydro-5H-thieno[2,3c]pyran-3-carboxylic acid tert-butyl ester (0.5 g, 1.21 mmol) was dissolved in dichloromethane (9 ml) and trifluoroacetic acid (3 ml) was added. The reaction mixture was stirred 64 hours at room temperature. The precipitate was filtered off and washed with diethyl ether and dried in vacuo at 50 °C for 4 hours, which afforded 180 mg (50 %) of the title compound as a solid.

.Mp.: 231 - 233 °C;

25 Calculated for C<sub>13</sub>H<sub>10</sub>F<sub>3</sub>NO<sub>8</sub>S;

C, 39.30 %; H, 2.56 %; N, 3.57 %. Found:

C, 39.30 %; H, 2.54 %; N, 3.53 %.

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5-(((Benzo[1,3]dioxol-5-ylmethyl)-amino)-methyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of oxalyl chloride (1 ml, 11.13 mmol) in dichloromethane (40 ml) cooled to -78 °C under an atmosphere of nitrogen was added dropwise a solution of dimethylsulfoxide (1.6 ml, 21.78 mmol) in dichloromethane (16 ml) during 5 min. After stirring for 15 min at -78 °C a solution of 2-(*tert*-butoxyoxalyl-amino)-5-hydroxymethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (2.0 g, 4.84 mmol) in dichloromethane (30 ml) was added dropwise and the resulting mixture was stirred for 0.5 hour at -78 °C. *N*,*N*-Diisopropylethylamine (4.2 ml, 24.18 mmol) was added and the reaction mixture allowed reaching room temperature at which time heptane (700 ml) was added. The mixture was filtered through anhydrous sodium sulfate and the solvent evaporated <u>in vacuo</u>. The residue (2.71 g) was purified on column chromatography using a mixture of ethyl acetate/heptane (1:4) as eluent which afforded 0.93 g (47 %) of 2-(*tert*-butoxyoxalyl-amino)-5-formyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil.

To a mixture of 2-(*tert*-butoxyoxalyl-amino)-5-formyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.46 g, 1.12 mmol) and piperonylamine (145  $\mu$ l, 1.12 mmol) in 1,2-dichloroethane (25 ml) was added sodium triacetoxyborohydride (0.35 g, 1.57 mmol) and the resulting mixture was stirred at room temperature for 1 hour. The mixture was washed with saturated aqueous sodium hydrogencarbonate (2 x 30 ml) and dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue (0.56 g) was purified on column chromatography using a mixture of ethyl acetate/heptane (1:1) as eluent followed by a mixture of 10% triethylamine in ethyl acetate/heptane (1:1) as eluent. Semi pure fractions were collected-and-the solvent-evaporated <u>in vacuo</u>. The residue (180 mg) was subjected to preparative TLC using a mixture of 10% triethylamine in ethyl acetate/ethanol (4:1) as eluent. The desired band was taken off and

extracted with methanol (400 ml) for 0.5 hour, filtered and the solvent evaporated in vacuo, which afforded 250 mg (> 100%, contains dichloromethane and silicagel) of 5-(((benzo[1,3]dioxol-5-ylmethyl)-amino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester as an oil. LC-MS: R<sub>t</sub> = 5.75 min, m/z: 547 [M+H]<sup>+</sup>.

LC-MS. Rt - 5.75 mm, m/z. 547 [M+n]

5-(((Benzo[1,3]dioxol-5-ylmethyl)-amino)methyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (250 mg) was dissolved in dichloromethane (9 ml) and trifluoroacetic acid (3 ml) was added. The reaction mixture was stirred 16 hours at room temperature. The volatiles were evaporated <u>in vacuo</u> and the residue trituated with a small portion of diethyl ether. The solid precipitate was filtered off and washed with diethyl ether and dried <u>in vacuo</u> at 50 °C for 16 hours, which afforded 160 mg of the <u>title compound</u> as a solid.

Calculated for C<sub>19</sub>H<sub>18</sub>N<sub>2</sub>O<sub>8</sub>S, 2xTFA, 3xH<sub>2</sub>O; C, 38.56 %; H, 3.66 %; N, 3.91 %. Found: C, 38.61 %; H, 3.90 %; N, 4.22 %.

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### **EXAMPLE 68**

5-((2-Methoxy-benzylamino)methyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-25 thieno[2,3-c]pyran-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in Example 66 using 2-(*tert*-butoxyoxalyl-amino)-5-formyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester and 2-methoxybenzylamine as starting material.

Calculated for C<sub>19</sub>H<sub>20</sub>N<sub>2</sub>O<sub>7</sub>S, 0.75xTFA; C, 48.67 %; H, 4.13 %; N, 5.54 %. Found: C, 48.61 %; H, 4.42 %; N, 5.35 %.

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### **EXAMPLE 69)**

5-((2-Benzo[1,3]dioxol-5-yl-acetylamino)methyl)-2-(oxalyl-amino)-4,7-20 dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

To a solution of 3,4-methylenedioxy phenylacetic acid (0.22 g, 1.09 mmol), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide, hydrochloride (0.27 g, 1.42 mmol) in acetonitrile (6 ml) was added triethylamine (0.46 ml, 3.27 mmol). The resultant mixture was allowed to stir at ambient temperature for 10 min. before 2-amino-5-aminomethyl-4,7-dihydro-5*H*-thieno-[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.311 g, 1.09 mmol) was added. The reaction mixture was allowed to stir at ambient temperature for 18 hours and then concentrated in vacuo. To the residue ethyl acetate and water were added and the layers separated. The organic layer was washed with hydrochloric acid (0.5M, (v/v)), saturated sodium bicarbonate (2 x 25 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The crude 2-amino-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)-methyl]-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester was used immediately in the next step.

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<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 6.78-6.69 (m, 3H), 5.97 (bs, 2H), 5.95 (s, 2H), 4.60-4.58 (m, 1H), 4.53 (s, 2H), 3.73 (ddd, 1H, J = 14 Hz, J = 7.6 Hz and J = 3.2 Hz), 3.65-3.59 (m, 1H), 3.49 (s, 2H), 3.11 (ddd, 1H, J = 12.4 Hz, J = 4 Hz and J = 4.4 Hz), 2.76 (dm, 1H), 2.44 (ddt, 1H, J = 19.6 Hz, J = 13.2 Hz and J = 2.4 Hz), 1.51 (s, 9H).

To a solution of the above crude 2-amino-5-((2-benzo[1,3]dioxol-5-ylacetylamino)-methyl]-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester (0.17 g, 0.38 mmol) in dichloromethane (5 ml) was added imidazol-1-yl-oxo-acetic acid tert-butyl ester (0.22 g, 1.14 mmol). The reaction mixture was stirred at room temperature for 18 hours, the volatiles evaporated in vacuo and the residue diluted with ethyl acetate. The organic layer was washed with hydrochloric acid (1% (v/v), 2 x 25 ml), saturated sodium bicarbonate (2 x 25 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered, concentrated in vacuo and the residue subjected to flash chromatography using a mixture of ethylacetate/hexanes (1:2) as eluent, which afforded 0.12 g (55 %) of 2-(tertbutoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)-methyl]-4,7dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  12.51 (bs, 1H), 6.78 (d, 1H, J = 8 Hz), 6.77 (d, 1H, J = 1.6 Hz), 6.71 (dd, 1H, J = 8.4 Hz and J = 1.6 Hz), 5.96 (s, 2H), 4.70 (m, 2H. J = 35 Hz. J = 15.2 Hz. J = 14.4 Hz and J = 2 Hz), 3.77 (ddd, 1H, J = 14.4 Hz and J =10.8 Hz, J = 7.6 Hz and J = 3.2 Hz), 3.67-3.62 (m, 1H), 3.50 (s, 2H), 3.15 (ddd, 1H, J = 12.8 Hz, J = 8.4 Hz and J = 4.4 Hz), 2.87 (dt, 1H, J = 16 Hz and J = 3 Hz), 2.57-2.50 (m, 1H), 1.61 (s, 9H), 1.57 (s, 9H); LC-MS: m/z: 575.0 [M+H]<sup>+</sup>

2-(*tert*-Butoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)-methyl]-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.12 g, 0.20 mmol) was dissolved in a 50% solution of trifluoroacetic acid/dichloromethane (2 ml). The reaction mixture was stirred at ambient temperature for 18 hours, concentrated in vacuo to 1/5 of the volume and

the precipitate filtered off and washed with dichloromethane (2x) affording 50 mg (50 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>)  $\delta$  12.32 (bs, 1H), 8.20 (t, 1H, J = 6.8 Hz), 6.81 (m, 2H), 6.70 (m, 1H), 5.95 (s, 2H), 4.80 (d, 1H, J = 19.6 Hz), 4.63 (d, 1H, J = 20 Hz), 3.65 (m, 1H), 3.34 (s, 2H), 3.30-3.20 (m, 3H), 2.87 (dm, 1H); LC-MS: m/z: 463.0 [M+H]<sup>+</sup>.

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### **EXAMPLE 70**

5-(((5-Methoxy-2-methyl-1*H*-indol-3-carbonyl)amino)methyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid

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To a solution of 5-methoxy-2-methyl indole-3-acetic acid (0.26 g, 1.18 mmol), and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide, hydrochloride (0.27 g, 1.4 mmol) in acetonitrile (10 ml) was added triethylamine (0.46 ml, 3.2 mmol). The reaction mixture was allowed to stir for 10 min at room temperature before compound 2-amino-5-aminomethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.307 g, 1.08 mmol) was added. The reaction mixture was allowed to stir for 18 hours and then concentrated in vacuo. Ethyl acetate and water were added and the layers separated. The organic layer was washed with hydrochloric acid (0.5M, (v/v)), saturated sodium bicarbonate (2 x 25 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo. The crude 2-amino-5-(((5-methoxy-2-methyl-1*H*-indol-3-carbonyl)amino)-methyl)-4,7-

dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester was used immediately in the next step.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 7.90 (bs, 1H), 7.19 (d, 1H, J = 8.8 Hz), 6.87 (d, 1H, J = 2.4 Hz), 6.79 (dd, 1H, J = 8.8 Hz and J = 2.4 Hz), 6.18 (m, 1H), 5.94 (s, 2H), 4.33 (m, 2H, J = 25 Hz, J = 14 Hz, J = 2.8 Hz and J = 1.6 Hz), 3.80 (s, 3H), 3.76 (ddd, 1H, J = 14 Hz, J = 8 Hz and J = 2.8 Hz), 3.65 (s, 3H), 3.53 (m, 1H), 2.99 (ddd, 1H, J = 13 Hz, J = 5.6 Hz and J = 4 Hz), 2.76 (dt, 1H, J = 16.8 Hz, J = 2.8 Hz), 2.42-2.40 (m, 1H), 2.38 (s, 3H), 1.51 (s, 9H).

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To a solution of the crude 2-amino-5-(((5-methoxy-2-methyl-1*H*-indol-3-carbonyl)amino)methyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.35 g, 0.72 mmol) in dichloromethane (5 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.42 g, 2.1 mmol).

- The reaction mixture was stirred at room temperature for 18 hours. The reaction mixture was concentrated in vacuo and the residue diluted with ethyl acetate. The organic layer was washed with hydrochloric acid (1% (v/v), 2 x 25 ml), saturated sodium bicarbonate (2 x 25 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered, concentrated in vacuo and the residue subjected to flash chromatography using a mixture of ethyl acetate/hexanes (1:1) as eluent, which afforded 0.24 (55 %) of 2-(tert-butoxyoxalyl-amino)-5-(((5-methoxy-2-methyl-1*H*-indol-3-carbonyl)amino)methyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic
- <sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 12.50 (bs, 1H), 7.92 (s, 1H), 7.20 (dd, 1H, J = 8.4 Hz and J = 0.4 Hz), 6.88 (d, 1H, J = 2.4 Hz), 6.80 (dd, 1H, J = 8.8 Hz and J = 2.4 Hz), 6.21 (m, 1H), 4.56 (dd, 1H, J = 14.8 Hz and J = 2.8 Hz), 4.44 (dt, 1H, J = 14.4 Hz and J = 2.8 Hz), 4.11 (q, 1H, J = 7.2 Hz), 3.81-3.75 (m, 1H), 3.79 (s', 3H), 3.66 (s, 2H), 3.58-3.54 (m, 1H), 3.01 (ddd, 1H, J = 14 Hz, J = 8.8 Hz and J = 4.4 Hz), 2.85 (dt, 1H, J = 16.8 Hz and J = 6 Hz), 2.52-2.45 (m, 1H), 2.38 (s, 3H), 1.60 (s, 9H), 1.57 (s, 9H);

acid tert-butyl ester as an oil.

LC-MS: m/z: 614.1 [M+H]<sup>+</sup>.

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2-(*tert*-Butoxyoxalyl-amino)-5-(((5-methoxy-2-methyl-1*H*-indol-3-carbonyl)amino)-methyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (0.24 g, 0.39 mmol) was dissolved in a 50 % solution of trifluoroacetic acid/dichloromethane (2 ml). The reaction mixture was stirred at ambient temperature for 18 hours, concentrated <u>in vacuo</u> to 1/5 of the volume and the precipitate filtered off. The filtrate was washed with dichloromethane (2x) and dried, which afforded 100 mg (50 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.31 (bs, 1H), 10.58 (s, 1H), 7.98 (t, 1H, J = 6.8 Hz), 7.08 (d, 1H, J = 11.2 Hz), 6.98 (d, 1H, J = 2.4 Hz), 6.58 (dd, 1H, J = 11.6 Hz/and J = 2.8 Hz), 5.75 (d, 1H, J = 0.8 Hz), 4.77 (d, 1H, J = 19.6 Hz), 4.58 (d, 1H, J = 20 Hz), 3.69 (s, 3H), 3.64-3.62(m, 1H), 3.43 (s, 2H), 3.31-3.20 (m, 1H), 2.92-2.84 (m, 1H), 2.52 (m, 1H-partially obscured by DMSO), 2.30 (s, 3H);

15 LC-MS: m/z: 500.1 [M-H].

20 <u>5-(1,3-Dioxo-5-propylcarbamoyl-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid</u>

In a 10-mL scintillating vial, a solution of 2-amino-5-aminomethyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (149 mg, 0.5 mmol) in *N*,*N*-dimethylformamide (4 mL) was treated with trimellitic anhydride (120 mg, 0.62 mmol) and stirred at 100 °C for 24 hours. The solution was then diluted with ethyl acetate (25 mL) and washed with 0.5N aqueous hydrogen chloride (25 mL) and brine (25 mL). The organic layer was dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo affording 229 mg (100 %) of 2-(2-amino-3-*tert*-butoxycarbonyl-4,7-dihydro-5*H*-thieno[2,3-

c]pyran-5-ylmethyl)-1,3-dioxo-2,3-dihydro-1*H*-isoindole-5-carboxylic acid as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>) δ 8.58 (s, 1H), 8.49 (d, 1H, J = 9 Hz), 8.00 (d, 1H, J = 10 Hz), 4.64-4.54 (m, 2H), 4.08-4.02 (m, 2H), 3.88-3.80 (m, 1H), 2.98-2.83 (m, 1H), 2.68-2.54 (m, 1H), 1.57 (s, 9H). HPLC (254.4 nm) R<sub>t</sub> = 3.98 min.

In a 250 mL round bottom flask, a solution of 2-(2-amino-3-tert-butoxycarbonyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-5-ylmethyl)-1,3-dioxo-2,3-dihydro-1*H*-isoindole-5-carboxylic acid (500 mg, 1.1 mmol) in dichloromethane (7 mL) was treated with a solution of imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (633 mg, 3.2 mmol) in dichloromethane (1.0 mL). After stirring for 4 hours at room temperature the reaction solution was dissolved in ethyl acetate (100 mL) and washed with distilled water (2 x 50 mL), 0.5 N aqueous hydrogen chloride (3 x 50 mL), and brine (50 mL). The organic layer was dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo to yield 370 mg (58 %) of 2-(2-(*tert*-butoxyoxalyl-amino)-3-*tert*-butoxycarbonyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-5-ylmethyl)-1,3-dioxo-2,3-dihydro-1*H*-isoindole-5-carboxylic acid as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>) δ 12.49 (s, 1H), 8.58 (s, 1H), 8.50 (d, 1H, J = 8 Hz), 8.00 (d, 1H, J = 8 Hz), 4.84-4.65 (m, 2H), 4.17-4.00 (m, 2H), 3.92-3.84 (m, 1H), 3.08-2.94 (m, 1H), 2.78-2.64 (m, 1H), 1.61 (s, 9H), 1.57 (s, 9H).

In a 50 mL round bottom flask, a solution of 2-(2-(*tert*-butoxyoxalyl-amino)-3-*tert*-butoxycarbonyl-4,7-dihydro-5*H*-thieno[2,3-c]pyran-5-ylmethyl)-1,3-dioxo-2,3-dihydro-1*H*-isoindole-5-carboxylic acid (208 mg, 0.36 mmol) in dichloromethane (5.0 mL) was treated with *N*,*N*-diisopropyl ethylamine (200 μL, 1.1 mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (84 mg, 0.44 mmol). The solution was allowed to stir for 50 minutes at room temperature before propylamine (30 μL, 0.36 mmol) was added dropwise. The solution was stirred for an additional 18 hours at room temperature. The volatiles were evaporated <u>in vacuo</u> and the

residue was purified by silica gel chromatography using a mixture of hexane/ethyl acetate (9:1) as eluent, which afforded 51 mg (23 %) of 2-(tert-butoxyoxalyl-amino)-5-(1,3-dioxo-5-propylcarbamoyl-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>) δ 12.48 (s, 1H), 8.24-8.16 (m, 2H), 7.93 (d, 1H, J = 8 Hz), 6.39 (t, 1H, J = 6 Hz), 4.18-4.63 (m, 2H), 4.10-3.96 (m, 2H), 3.92-3.78 (m, 1H), 3.47 (q, 2H, J = 7 Hz), 2.99 (d, 1H, J = 17), 2.76-2.60 (m, 1H), 1.68 (q, 2H, J = 7 Hz), 1.61 (s, 9H), 1.57 (s, 9H), 1.01 (t, 3H, J = 7 Hz).

In a 25 mL round bottom flask 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-5-propylcarbamoyl-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid *tert*-butyl ester (40 mg, 0.07 mmol) was dissolved in 20 % trifluoroacetic acid in dichloromethane (4 mL). The solution was left open to the atmosphere without stirring. After 24 hours the precipitate was filtered off and washed with diethyl ether, affording 32 mg (90 %) of the <u>title compound</u> as a solid.

<sup>1</sup>H-NMR (300 MHz, DMSO-d<sub>6</sub>) δ12.32 (s, 1H), 8.81 (s, 1H), 8.58 (s, 1H), 8.00 (s, 1H), 4.90-4.48 (m partially obscured by water, 2H), 4.00-3.64 (m partially obscured by water, 3H), 3.36-3.16 (m partially obscured by water, 2H), 3.13-2.90 (d partially obscured by water, 1H), 2.69-2.53 (m partially obscured by DMSO, 1H), 1.69-1.38 (m, 2H), 1.00-0.74 (m, 3H).

25 HPLC (254.4 nm) R<sub>t</sub> = 3.09 min.MS (APCI) m/z: 515.4 [M-H].

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7-(4-Hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

2-(*tert*-Butoxyoxalyl-amino)-7-(4-benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (50 mg, 0.067 mmol) was dissolved in a mixture of ethyl acetate/ethanol (3 mL, 1:1). Palladium on activated carbon (10%, 10 mg) was added and the solution degassed and stirred under hydrogen (1 atm) for 72 hours. The mixture was filtered through celite and the filter cake washed with hot ethyl acetate. The filtrate was concentrated <u>in vacuo</u> and the residue purified by silica gel chromatography (10% ethyl acetate/dichloromethane) to obtain 42 mg (95%) of 2-(*tert*-butoxyoxalyl-amino)-7-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-

an oil.

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<sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>) δ 12.59-12.53 (2s, 1H), 7.64-7.53 (m, 1H), 7.42-7.36 (m, 1H), 7.19-7.11 (m, 1H), 5.58-5.37 (m, 1H), 4.37-4.00 (m, 2H), 3.86-3.78 (m, 1H), 3.32-3.18 (m, 1H), 2.99-2.94 (m, 1H), 2.84-2.69 (m, 1H), 1.62-1.59 (3s, 18H), 1.17-1.11 (2s, 9H);

tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester as

20 LC-MS:  $R_t = 4.55 \text{ min, m/z}$ : 658 [M+H]<sup>+</sup>,

2-(*tert*-Butoxyoxalyl-amino)-7-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di*tert*-butyl ester (42 mg, 0.064 mmol) was dissolved in a solution of 50% trifluoroacetic acid/methylene chloride (3 mL). The reaction was stirred at ambient temperature for 7 hours, concentrated <u>in vacuo</u> and evaporated from dichloromethane (10 ml) three times. The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 29 mg (81 %) of the title compound as a solid trifluoroacetate.

<sup>1</sup>H-NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.32 (bs, 1H), 11.26 (s, 1H), 9.30 (bs, 2H), 7.64 (t, 1H, J = 7 Hz), 7.33 (d, 1H, J = 7 Hz), 7.25 (d, 1H, J = 7 Hz), 4.84 (s, 1H), 4.06-3.96 (m, 2H), 3.56 (m, 2H), 3.05 (bs, 2H), LC-MS:  $R_t$  = 1.26 min, m/z:  $[M+H]^+$ ,

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### **EXAMPLE 73**

5-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

Acetyl chloride (5.4 ml, 5.96 g, 76 mmol) was added dropwise to methanol (15 ml) at 0 °C in a sealed 50 ml round-bottom flask. This solution was allowed to warm to room temperature for 1 hour while stirring. To this solution 3-hydroxy-2-methyl-benzoic acid (519 mg, 3.4 mmol) was added and the solution was stirred at room temperature for 42 hours. The reaction was quenched with saturated aqueous sodium bicarbonate and solid sodium bicarbonate. The volatiles were removed in vacuo and the basic aqueous solution was then extracted with dichloromethane (4 x 40 ml). The combined organic extracts were dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo affording 493 mg (87 %) of 3-hydroxy-2-methyl-benzoic acid methyl ester as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  7.43 (d, 1H, J = 9 Hz), 7.12 (t, 1H, J = 8 Hz), 6.95 (d, 1H, J = 8 Hz), 5.05 (bs, 1H), 3.90 (s, 3H), 2.47 (s, 3H).

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To a solution of the above methyl ester (256 mg, 1.54 mmol) and N,N-diisopropylethylamine (530  $\mu$ l, 3.0 mmol) in dichloromethane (8 ml) at 0 °C methyloxymethyl chloride (175  $\mu$ l, 2.3 mmol) was added dropwise. The solution was allowed slowly to warm to room temperature and stired for 24 hours. The solution was diluted with dichloromethane (12 ml), washed with

water (20 ml), brine (20 ml), dried (MgSO<sub>4</sub>), filtered, and concentrated <u>in vacuo</u>. The resulting oil was purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (4:1) as eluent, which afforded 269 mg (85 %) of 3-methoxymethoxy-2-methyl-benzoic acid methyl ester as an oil.  $^{1}$ H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  7.48 (d, 1H, J = 8 Hz), 7.24-7.15 (m, 2H), 5.22 (s, 2H), 3.90 (s, 3H), 3.50 (s, 3H), 2.47 (s, 3H).

In a 25 ml round-bottom flask, *N*-bromosuccinimide (236 mg, 1.3 mmol) and azobis(cyclohexanecarbonitrile) (33 mg, 0.14 mmol) were added to a solution of 3-methoxymethoxy-2-methyl-benzoic acid methyl ester (265 mg, 1.26 mmol) in carbon tetrachloride (6.5 ml). The reaction was heated to reflux with stirring for 3.5 hours. The volatiles were removed <u>in vacuo</u> and the residue purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (9:1) as eluent, which afforded 364 mg (100 %) of 2-bromomethyl-3-methoxymethoxy-benzoic acid methyl ester as a solid. 

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  7.55 (dd, 1H, J = 6,3 Hz), 7.29 (d, 2H, J = 3 Hz), 5.27 (s, 2H), 5.05 (s, 2H), 3.91 (s, 3H), 3.50 (s, 3H).

In a 100 ml round-bottom flask, 2-amino-5-aminomethyl-6-(4-methoxy-20 benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (298 mg, 0.74 mmol) and N,N-diisopropylethylamine (195 µl, 1.12 mmol) were dissolved in acetonitrile (40 ml). 2-Bromomethyl-3methoxymethoxy-benzoic acid methyl ester (193 mg, 0.67 mmol) in acetonitrile (5 ml) was slowly added to the amine solution via gastight 25 syringe over 24 hours, followed by stirring at room temperature for an additional 36 hours. The solution was concentrated in vacuo, the residue redissolved in ethyl acetate (25 ml), and washed with saturated aqueous sodium bicarbonate (25 ml) and brine (25 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (1:1) as eluent, which afforded 345 mg (81 %) of 2-amino-6-(4methoxy-benzyl)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-

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ylmethyl)-4,5,6,7-tetrahydro[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  7.67 (d, 1H, J = 8 Hz), 7.57-7.38 (m, 5H), 7.14 (d, 2H, J = 8 Hz), 6.96 (m, 2H), 6.77 (d, 2H, J = 9 Hz), 6.20 (d, 2H, J = 6 Hz), 5.96 (s, 2H), 4.69-2.58 (m, 17H), 1.55 (s, 9H).

In a 50 ml round-bottom flask a solution of 2-amino-6-(4-methoxy-benzyl)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7tetrahydro[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (338 mg, 0.58 mmol) in dichloromethane (20 ml) was treated with imidazol-1-vl-oxoacetic acid tert-butyl ester (575 mg, 2.9 mmol). After stirring for 18 hours at room temperature, the mixture was concentrated to dryness in vacuo. The residue was purified by silica gel chromatography using a mixture of hexanes/ethyl acetate (1:1) as eluent, which afforded 310 mg (75 %) of 2-(tert-Butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(4-methoxymethoxy-1oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid. <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  12.57 (s, 1H), 7.53 (d, 1H, J = 8 Hz), 7.43 (t, 1H. J = 8 Hz), 7.26 (d, 1H, J = 8 Hz), 7.13 (d, 2H, J = 9 Hz), 6.78 (d, 2H, J= 9 Hz), 5.28 (s, 2H), 4.47 (q, 2H, J = 18 Hz), 4.02-3.44 (m, 11H), 2.97 (dd, 1H, J = 18 Hz and J = 5 Hz), 2.76 (dd, 1H, J = 17 Hz and J = 5 Hz),1.63 (s, 9H), 1.59 (s, 9H).

10 % Pd/C (145 mg, 50 % by weight) was added to a mixture of 2-(*tert*-butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (283 mg, 0.40 mmol) in 10 % formic acid and methanol (10 ml). After stirring at room temperature for 18 hours, more Pd/C (141 mg, 50 % by weight) was added to the reaction mixture.

After stirring at room temperature for an additional 20 hours, the catalyst was removed via fitration through celite. Fresh Pd/C (255 mg) and ammonium formate (1.0 g) were added to the residue (253 mg, 0.36 mmol) dissolved in 10 % formic acid in methanol (10 ml). The solution was

heated to 40 °C for 48 hours. Catalyst was removed via filtration through celite and liberal washing with methanol. Purification by chromatotron (ethyl acetate/triethylamine (99:1)) afforded 63 mg (27 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester **A** and 46 mg (19 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-methyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester **B**.

- A: <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): δ 12.54 (s, 1H), 7.50 (d, 1H, J = 8 Hz), 7.41 (t, 1H, J = 8 Hz), 7.25 (d, 1H, J = 8 Hz), 5.27 (s, 2H), 4.52 (dd, 2H, J = 30 Hz and J = 19 Hz), 4.08-3.90 (m, 2H), 3.86-3.67 (m, 2H), 3.51 (s, 3H), 3.27 (m, 1H), 2.99 (dd, 1H, J = 18 Hz and J = 4 Hz), 2.53 (dd, 1H, J = 18 Hz and J = 11 Hz), 1.61 (s, 9H), 1.53 (s, 9H).
- 15 LC-MS (APCl<sup>+</sup>) m/z: 588 [M+H]<sup>+</sup>; R<sub>t</sub> = 1.32 min.

B: <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): δ 12.56 (s, 1H), 7.50 (d, 1H, J = 7 Hz), 7.41 (t, 1H, J = 8 Hz), 7.25 (d, 1H, J = 8 Hz), 5.27 (s, 2H), 4.50 (dd, J = 28 Hz and J = 18 Hz), 3.93-3.68 (m, 4H), 3.51 (s, 1H), 3.51 (s, 3H), 3.31 (m, 1H),
2.88 (dd, 1H, J = 18 Hz and J = 4 Hz), 2.68 (dd, 1H, J = 19 Hz and J = 9 Hz), 2.46 (s, 3H), 1.61 (s, 9H), 1.54 (s, 9H).
LC-MS (APCl<sup>+</sup>) m/z: 602 [M+H]<sup>+</sup>; R<sub>t</sub> = 1.35 min.

2-(tert-Butoxyoxalyl-amino)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester A (63 mg, 0.11 mmol) was dissolved in 30 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring. After 24 hours the precipitate was filtered off and washed with diethyl ether, affording 57 mg (90 %) of the title
 compound as a solid trifluoroacetate.

<sup>1</sup>H-NMR (300 MHz, DMSO-d<sub>6</sub>): δ 12.30 (s, 1H), 10.17 (s, 1H), 9.23 (s, 2H, J = 5 Hz and J = 7 Hz), 7.34 (t, 1H, J = 6 Hz), 7.19 (d, 1H, J = 5 Hz), 7.03 (d, 1H, J = 6 Hz), 5.76 (s, 2H), 4.53 (d, 1H, J = 13 Hz), 4.43-4.22 (m, 3H),

4.07 (m, 1H), 3.91 (m, 1H), 3.70 (m, 1H), 3.10 (m, 1H), 2.82 (dd, 1H, J = 14 Hz and J = 8 Hz).

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### **EXAMPLE 74**

5-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-methyl-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

- The above 2-(*tert*-butoxyoxalyl-amino)-5-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-methyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester **B** (46 mg, 0.08 mmol) was dissolved in 30 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring. After 24 hours
- the precipitate was filtered off and washed with diethyl ether, affording 41 mg (90 %) of the <u>title compound</u> as a solid trifluoroacetate.

  <sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>): δ 12.39 (s, 1H), 10.19 (s, 1H), 10.10 (s, 1H), 7.32 (t, 1H, J = 7.6 Hz), 7.17 (d, 1H, J = 7.2 Hz), 7.02 (t, 1H, J = 7.2 Hz), 4.55 (d, 2H, J = 15 Hz), 4.0-4.5 (m, 4H), 2.95-3.70 (m, 5H), 2.85 (s, 3H).
- 20 LC-MS (APCI<sup>+</sup>) m/z: 446 [M+H]<sup>+</sup>; R<sub>t</sub> = 1.02 min.

### **EXAMPLE 75**

25 5=((1,1-Dioxo-1*H*-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

Saccharin (8.8 g, 48 mmol) and phosphorous pentachloride (15 g, 72 mmol) were added neat to a round bottom flask equipped with a short path distillation column. The mixture was heated to 175 °C. After approximately 0.5 hour, phosphorous oxychloride slowly distilled off.

Upon completion of the reaction, the mixture was cooled and the resultant solid recrystallized from benzene affording 3.6 g (37 %) of 3-chlorobenzo[d]isothiazole 1,1-dioxide as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.92 (d, 1H, J = 6.9 Hz), 7.8 (m, 3H).

To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-10 tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (155 mg, 0.384 mmol) and triethylamine (59 µl, 0.423 mmol) in dichloromethane (2 ml) at 0 °C, was added a solution of 3-chloro-benzo[d]isothiazole 1,1dioxide (85.2 mg, 0.423 mmol) in dichloromethane (2 ml). The reaction mixture was stirred at 0 °C for 1hour. The reaction was judged complete 15 by tlc (dichloromethane/ethyl acetate (1:1)). The reaction mixture was washed with water (3 x 20 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The crude residue was subjected to flash chromatography using a gradient from 100 % dichloromethane to dichloromethane/ethyl acetate (80/20) as eluent, which afforded 200 mg 20 (92 %) of 2-amino-5-((1,1-dioxo-1H-benzo[d]isothiazol-3-ylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as a foam.

<sup>1</sup>H-NMR (CD<sub>3</sub>OD): δ 7.99 (m, 1H), 7.87 (m, 1H), 7.79 (m, 2H), 7.19 (d, 2H, J = 8.4 Hz), 6.75 (d, 2H, J = 8.7 Hz), 3.88-3.79 (m, 2H), 3.75-3.59 (m, 3H), 3.69 (s, 3H), 3.52-3.46 (m, 2H), 2.84 (dd, 1H, J = 15.3 Hz and J = 5.4 Hz), 2.68 (dd, J = 18 Hz and J = 4.5 Hz), 1.46 (s, 9H). *LC-MS*: R<sub>t</sub> = 2.83, m/z: 569 [M+H]<sup>+</sup>

To a solution of 2-amino-5-((1,1-dioxo-1H-benzo[d]isothiazol-3-ylamino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (129 mg, 0.227 mmol) in tetrahydrofuran (3 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (1.1 ml, 1.1

mmol, 1 M in tetrahydrofuran). The reaction mixture was stirred at room temperature for 18 hours. The reaction mixture was concentrated *in vacuo* and the residue subjected to flash chromtography using a mixture of ethyl acetate/dichloromethane (10:90) as eluent, which afforded 142 mg (90%) of 2-(*tert*-butoxyoxalyl-amino)-5-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.  $^{1}$ H-NMR (CDCl<sub>3</sub>):  $\delta$  7.92 (d, 1H, J = 6.3 Hz), 7.73 (m, 2H), 7.56 (d, 1H, J = 5.7 Hz), 7.20 (d, 2H, J = 6.3 Hz), 7.05 (bs, 1H), 6.87 (d, 2H, J = 6.6 Hz), 3.91 (m, 2H), 3.82-3.72 (m, 2H), 3.79 (s, 3H), 3.61-3.49 (m, 2H), 3.44 (m, 1H), 3.11 (dd, 1H, J = 15 Hz and J = 3.6 Hz), 2.72 (dd, 1H, J = 12 Hz and J = 4.2 Hz), 1.63 (s, 18H); LC-MS:  $R_t$ =3.48, m/z: 697 [M+H]<sup>†</sup>

- 2-(tert-Butoxyoxalyl-amino)-5-((1,1-dioxo-1H-benzo[d]isothiazol-3-15 ylamino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid *tert*-butyl ester (120 mg, 0.172 mmol) was dissolved in a mixture of ethanol (4 ml) and formic acid (0.5 ml). 10 % Pd-C (20 mg) was added and the reaction mixture stirred at ambient <sub>7</sub> 20 temperature for 4 days (after the second day, 150 mg of additional 10 % Pd-C was added). The reaction mixture was filtered through celite and the celite washed with dichloromethane. The organic fractions were combined and concentrated in vacuo. The resultant oil was subjected to preparative thin layer chromatography (dichloromethane/methanol (95:5)), which 25 afforded 17 mg (17 %) of 2-(tert-butoxyoxalyl-amino)-5-((1,1-dioxo-1Hbenzo[d]isothiazol-3-ylamino)-methyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid. <sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 7.91 (m, 1H), 7.72 (m, 3H), 7.34 (bs, 1H), 4.16-4.08 (m, 1H), 4.07 (dd, 2H, J = 36.3 Hz and J = 8.7 Hz), 3.38-3.30 (m, 1H),30 3.22-3.06 (m, 2H), 2.51 (dd, 1H, J = 16.8 Hz and J = 9.9 Hz), 1.61 (s, 18H).
  - 2-(*tert*-Butoxyoxalyl-amino)-5-((1,1-dioxo-1H-benzo[d]isothiazol-3-ylamino)-methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

tert-butyl ester (15 mg, 0.026 mmol) was dissolved in a solution of 50 % trifluoroacetic acid/dichloromethane (3 ml). The reaction mixture was stirred at ambient temperature for 18 hours, concentrated <u>in vacuo</u> and reevaporated from acetonitrile (2x). The residue was washed with dichloromethane and dried <u>in vacuo</u> to give 16 mg (90 %) of the <u>title</u>

<sup>1</sup>H-NMR (CD<sub>3</sub>OD):  $\delta$  7.98 (d, 1H, J = 7.2 Hz), 7.92 (d, 1H, J = 6.6 Hz), 7.83 (m, 2H), 4.51-4.39 (m, 2H), 4.11-4.08 (m, 1H), 3.97-3.91 (m, 2H), 3.53-3.47 (m, 1H), 3.16-3.10 (m, 1H).

compound as a solid trifluoroacetate.

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## **EXAMPLE 76**

7-((1,1-Dioxo-1H-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino) 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid 3-Chloro-benzo[d]isothiazole-1,1-dioxide (160 mg, 0.79 mmol) and diisopropylethylamine (150 µl, 0.86 mmol) were dissolved in dichloromethane (7 ml) at 0 °C. 2-Amino-7-aminomethyl-6-(4-methoxybenzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (284 mg, 0.70 mmol) was added and the mixture was stirred for 15 20 minutes at 0 °C, diluted with dichloromethane (10 ml) and washed with water (20 ml) and brine (20 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated in vacuo. The residue was purified by silica gel chromatography using a gradient of hexanes/ethyl acetate (1:1) to pure ethyl acetate as eluent, which afforded 309 mg (77 %) of 2-amino-25 7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)methyl)-6-(4-methoxybenzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an foam.

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<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>):  $\delta$  7.89 (d, 1H, J = 8 Hz), 7.77-7.63 (m, 2H), 7.37 (d, 1H, J = 7 Hz), 7.25 (d, 2H, J = 10 Hz), 6.82 (d, 2H, J = 8 Hz), 6.62 (bs, 1H), 6.08 (s, 2H), 3.91 (m, 1H), 3.71 (s, 3H), 3.49-2.65 (m, 8H), 1.59 (s, 9H).

5 LC-MS (APCI<sup>+</sup>) m/z: 569 [M+H]<sup>+</sup>, [M+Na] 591;  $R_t = 2.85$  min.

2-Amino-7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (102 mg, 0.18 mmol) in dichloromethane (10 ml) was treated with imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (85 mg, 0.43 mmol). After stirring for 18 hours at room temperature, the reaction solution was concentrated to dryness <u>in vacuo</u>. The residue was purified by silica gel chromatography using a gradient of hexanes/ethyl acetate (1:1) to pure ethyl acetate as gradient, which afforded 98 mg (78 %) of 2-(*tert*-butoxyoxalyl-amino)-7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): δ 12.57 (s, 1H), 7.89 (d, 1H, J = 8 Hz), 7.77-7.63 (m, 2H), 7.39 (d, 1H, J = 7 Hz), 7.25 (d, 2H, J = 9 Hz), 6.84 (d, 2H, J = 9 Hz), 6.64 (bs, 1H), 3.99-2.76 (m, 12H), 1.64 (s, 9H), 1.63 (s, 9H). 10 % Pd/C (100 mg) was added to a mixture of 2-(*tert*-butoxyoxalyl-amino)-7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (98 mg, 0.14 mmol) in 10 % formic acid in methanol (10 ml). After stirring at room temperature for 48 hours, the catalyst was removed via filtration through celite and liberal washing with methanol. The volatiles were removed <u>in vacuo</u> and the residue purified by chromatotron (ethyl acetate/triethylamine, 99:1), which afforded 32 mg (40 %) of 2-(*tert*-butoxyoxalyl-amino)-7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)-methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): δ 12.48 (s, 1H), 10.21-9.15 (m, 2H), 8.49-7.42 (m, 3H), 5.62-5.00 (bs, 1H), 4.53-2.87 (m, 8H), 1.61 (s, 18H).

HPLC (254.4 nm)  $R_t = 3.67$  minutes.

2-(*tert*-Butoxyoxalyl-amino)-7-((1,1-dioxo-1*H*-benzo[d]isothiazol-3-ylamino)-methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (32 mg) was dissolved in a mixture of 30 % trifluoroacetic acid in dichloromethane (4 ml). The solution was left open to the atmosphere without stirring. After 24 hours the precipitate was filtered off and washed with diethyl ether, affording 29 mg (90 %) of the <u>title</u> <u>compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (300 MHz, DMSO-d<sub>6</sub>): δ 12.36 (s, 1H), 9.92 (bs, 1H), 9.73 (bs, 1H), 9.38 (bs, 1H), 8.20 (m, 1H), 8.05 (m, 1H), 7.89 (m, 2H), 4.95 (s, 1H), 4.12-3.00 (m partially obscured by water, 8H). LC-MS (APCI<sup>+</sup>) m/z: 466 [M+H]<sup>+</sup>; R<sub>t</sub> = 0.66 min.

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### **EXAMPLE 77**

5-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

2-Methoxy-6-methylbenzoic acid ethyl ester (500 mg, 2.67 mmol), N-bromosuccinimide (483.8 mg, 2.72 mmol) and 2,2'-azobis(2-methyl-propionitrile) (30.2 mg, 0.123 mmol) in carbon tetrachloride (10 ml) were heated to reflux. After 18 hours, the reaction mixture was evaporated to dryness in vacuo. The residue was dissolved in dichloromethane (100 ml) and washed with water (2 x 50 ml). The organic layer was dried (MgSO<sub>4</sub>),

filtered and the solvent evaporated in vacuo. The residue (702 mg) was purified by column chromatography using a mixture of hexanes/dichloromethane (1:1) as eluent, which afforded 573 mg (85 %) of 6-bromomethyl-2-methoxy-benzoic acid ethyl ester as an oil.  $^{1}$ H-NMR (CDCl<sub>3</sub>):  $\delta$  7.37 (t, 1H, J = 8.4 Hz), 7.01 (d, 1H, J = 8.1 Hz), 6.90 (d, 1H, J = 8.4 Hz), 4.54 (s, 2H), 4.45 (q, 2H, J = 7.2 Hz), 3.82 (s, 3H), 1.42 (t, 3H, J = 9 Hz).

6-Bromomethyl-2-methoxy-benzoic acid ethyl ester (71.1 mg, 0.260 mmol) dissolved in acetonitrile (5 ml) and diisopropylethylamine (453 µl, 2.60 mmol) was stirred at room temperature. To this mixture 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3carboxylic acid tert-butyl ester (200 mg, 0.52 mmol) dissolved in acetonitrile (5 ml) was added syringe pump (0.2 ml/min.). Once addition was complete, the reaction mixture was allowed to stir for 2 hours. The reaction mixture was concentrated in vacuo, and the residue diluted with ethylacetate (50 ml). The organic layer was washed with saturated sodium bicarbonate (2 x 25 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo. The residue (308 mg) was subjected to column chromatography using a gradient of hexane/ethylacetate (95:5) to (50:50) and then dichloromethane/ethyl acetate (95:5) as eluents, which afforded 106 mg (75 %) of 2-amino-6-(4-methoxy-benzyl)-5-(7-methoxy-1-oxo-1.3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.48 (t. 1H, J = 7.5 Hz), 7.12 (d, 2H, J = 8.4 Hz), 7.01 (d. 1H, J = 7.5 Hz), 6.91 (d. 1H, J = 8.4 Hz), 6.76 (d. 2H, J = 7.8 Hz), 5.95

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To a solution of 2-amino-6-(4-methoxy-benzyl)-5-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (105 mg, 0.192 mmol) in tetrahydrofuran (3

(bs, 2H), 4.37 (s, 2H), 4.05 (m, 1H), 3.97 (s, 3H), 3.88-3.78 (m, 2H), 3.81 (s, 3H), 3.71-3.39 (m, 4H), 2.90 (dd, 1H, J = 18 Hz and J = 5.4 Hz), 2.62

(dd, 1H, J = 18 Hz and J = 5.4 Hz), 1.53 (s, 9H).

ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.534 ml, 0.534 mmol, 1 M in tetrahydrofuran). The reaction mixture was stirred at room temperature for 18 hours. The reaction mixture concentrated <u>in vacuo</u> and the residue subjected to flash chromtography using a mixture of ethyl acetate/dichloromethane (10:90) as eluent, which afforded 85 mg (66 %) of 2-(*tert*-butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 7.47 (t, 1H, J = 5.7 Hz), 7.10 (d, 2H, J = 6 Hz), 6.99 (d, 1H, J = 5.7 Hz), 6.90 (d, 1H, J = 6.3 Hz), 6.76 (d, 2H, J = 6.3 Hz), 4.37 (q, 2H, J = 11.4 Hz), 3.99-3.92 (m, 1H), 3.97 (s, 3H), 3.79-3.76 (m, 2H), 3.77 (s, 3H), 3.66 (d, 1H, J = 12.6 Hz), 3.58-3.50 (m, 3H), 2.95 (dd, 1H, J = 13.5 Hz and J = 3.6 Hz), 2.70 (dd, 1H, J = 13.5 Hz and J = 3.6 Hz), 1.61 (d, 9H), 1.57 (s, 9H).

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2-(*tert*-Butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (66 mg, 0.12 mmol) was dissolved in ethanol (2 ml) and formic acid (0.3 ml). 10 % Pd-C (15 mg) was added and the reaction mixture stirred at room temperature for 3 days. TLC (hexane/ethyl acetate (1/1)) indicated reaction complete. The reaction mixture was filtered through celite and the celite washed with dichloromethane. The organic fractions were combined and subjected to preparative thin layer chromatography (hexane/ethyl acetate (1/1) to yield 14.7 mg (22 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.48 (t, 1H, J = 7.5 Hz), 7.01 (d, 1H, J = 7.2 Hz), 6.90 (d, 1H, J = 8.4 Hz), 5.50 (d, 2H, J = 6.6 Hz), 4.04-3.90 (m, 1H), 3.97 (s, 3H), 3.24 (m, 1H), 3.01-2.95 (m, 1H), 2.57-2.43 (m, 2H), 1.62 (s, 9H), 1.57 (s, 9H).

2-(*tert*-Butoxyoxalyl-amino)-5-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-

butyl ester (14.7 mg, 0.026 mmol) was dissolved in a solution of 50% trifluoroacetic acid/dichloromethane (2 ml). The reaction mixture was stirred at ambient temperature for 18 hours, concentrated <u>in vacuo</u> and reevaporated from acetonitrile (2x). The resulting precipitate was washed with dichloromethane and dried <u>in vacuo</u> to give 13 mg (89 %) of the <u>title</u> compound as a solid trifluoroacetate.

<sup>1</sup>H-NMR (CD<sub>3</sub>OD):  $\delta$  7.56 (t, 1H, J = 8.1 Hz), 7.13 (d, 1H, J = 7.2 Hz), 7.01 (d, 1H, J = 8.1 Hz), 4.87-4.44 (m, 4H), 4.15 (m, 1H), 3.90 (s, 3H), 3.88-3.79 (m, 1H), 3.43 (m, 1H), 2.98 (m, 2H);

10 LC-MS:  $R_t = 0.71$ , m/z: 446 [M+H]<sup>+</sup>.

### **EXAMPLE 78**

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5-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of 2-hydroxy-6-methyl-benzoic acid ethyl ester (5.00 g, 27.8 mmol) and t-butyl-di-methylsilyl chloride (6.27 g, 41.6 mmol) in dichloromethane (100 ml) was added diisopropyl ethylamine. The solution was stirred at 50 °C for 24 hours, washed with water, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo, which afforded 7.6 g (93 %) of 2-(*tert*-butyl-dimethyl-silanyloxy)-6-methyl-benzoic acid ethyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.13 (t, 1H, J = 7.5 Hz), 6.78 (d, 1H, J = 7.5 Hz), 6.67 (d, 1H, J = 7.5 Hz), 4.35 (q, 2H, J = 7.2 Hz), 2.29 (s, 3H), 1.38 (t, 3H, J = 7.2 Hz), 0.97 (s, 9H), 0.23 (s, 6H).

2-(tert-Butyl-dimethyl-silanyloxy)-6-methyl-benzoic acid ethyl ester (7.6 g,
 25.8 mmol), N-bromosuccinimide (4.82 g, 27.1 mmol) and
 azobis(cyclohexanecarbonitrile) (0.32 g, 1.3 mmol) were dissolved in

tetrachlormethane (130 ml). The solution was stirred at room temperature for 60 hours. The solvent was evaporated in vacuo and the residue was chromatographed on silica gel column using a gradient of 1-2% ethyl acetate/hexane as eluent, which afforded 8.0 g (83 %) of 6-bromomethyl-2-(*tert*-butyl-dimethyl-silanyloxy)-benzoic acid ethyl ester as an oil.  $^{1}$ H-NMR (CDCl<sub>3</sub>):  $\delta$  7.21 (t, 1H, J = 8.4 Hz), 7.00 (d, 1H, J = 8.4 Hz), 6.81 (d, 1H, J = 8.4 Hz), 4.51 (s, 2H), 4.40 (q, 2H, J = 7.2 Hz), 1.42 (t, 3H, J = 7.2 Hz), 0.98 (s, 9H), 0.23 (s, 6H).

- To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-10 tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (3.00 g, 7.45 mmol) and diisopropyl ethylamine (1.93 ml, 11.2 mmol) in acetonitrile at room temperature was added a solution of 6-bromomethyl-2-(tert-butyldimethyl-silanyloxy)-benzoic acid ethyl ester (2.78 g, 7.45 mmol) in acetonitril over 48 hours. The solution was stirred for 12 hours after the 15 addition was complete. The volatiles were evaporated in vacuo and the residue was taken into ethyl acetate (50 ml) and washed with water, 1 N hydrochloric acid, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was chromatographed on silica gel column eluted with a mixture of 20 % ethyl acetate/Hexane, which -20afforded 3.2 g (66 %) of 2-amino-5-(7-(tert-butyl-dimethyl-silanyloxy)-1oxo-1,3-dihydro-isoindol-2-ylmethyl]-6-(4-methoxy-benzyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil. <sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.36 (t, 1H, J = 8.0 Hz), 7.11 (d, 2H, J = 8.8 Hz), 6.99 (d. 1H, J = 8.0 Hz), 6.82 (d. 1H, J = 8.0 Hz), 6.76 (d. 2H, J = 8.8 Hz), 5.94 25 (s, 2H), 4.48 (d, 1H, J = 16.8 Hz), 4.33 (d, 1H, J = 16.8 Hz), 3.90-3.45 (m, 7H), 3.78 (s, 3H), 2.95 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 2.72 (dd, 1H, J = 17 Hz and J = 5.6 Hz), 1.52 (s, 9H), 1.05 (s, 9H), 0.26 (s, 6H).
- To a stirred solution of 2-amino-5-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (2.37 g, 3.64 mmol) in tetrahydrofuran (50 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-

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butyl ester (2.14 mg, 10.9 mmol) in tetrahydrofuran (10 ml). The mixture was stirred at room temperature for 24 hours. The solvent was removed in vacuo. The residue was taken into ethyl acetate (100 ml). The solution was washed with 0.5 N hydrochloric acid solution (2 x 20 ml), saturated sodium bicarbonate (2 x 20 ml) and brine (20 ml), dried (MgSO<sub>4</sub>), filtered and the solvent removed in vacuo. The residue was chromatographed using a gradient of 10-20 % ethyl acetate/Hexane as eluent, which afforded 2.40 g (92 %) of 2-(tert-butoxyoxalyl-amino)-5-(7-(tert-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.59 (s, 1H), 7.37 (t, 1H, J = 8.0 Hz), 7.10 (d, 2H, J = 8.8 Hz), 7.00 (d, 1H, J = 8.0 Hz), 6.83 (d, 1H, J = 8.0 Hz), 6.77 (d, 2H, J = 8.8 Hz), 4.50 (d, 1H, J = 16.8 Hz), 4.34 (d, 1H, J = 16.8 Hz), 3.90-3.45 (m, 7H), 3.77 (s, 3H), 2.95 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 2.72 (dd, 1H, J = 18 and J = 5.6 Hz), 1.61 (s, 9H), 1.58 (s, 9H), 1.06 (s, 9H), 0.26 (s, 6H).

ester as a solid.

To a solution of 2-(*tert*-butoxyoxalyl-amino)-5-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (2.40 g, 3.34 mmol) in 10 % formic acid/methanol (50 ml) at room temperature under nitrogen was added 10 % Pd/C (1.2 g). The mixture was stirred for 48 hours. The Pd/C was filtered off and the filtrate was evaporated <u>in vacuo</u>. The residue was dissolved in dichloromethane (10 ml). The resulting solution was poured into hexane. The precipitate was filtered off and dried <u>in vacuo</u> affording 1.3 g (61 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.45 (s, 1H), 8.05 (s, 1H), 7.39 (t, 1H, J = 8.0 Hz), 7.00 (d, 1H, J = 8.0 Hz), 6.83 (d, 1H, J = 8.0 Hz), 4.50 (d, 1H, J = 16.8 Hz), 4.45 (q, 2H, J = 17 Hz), 4.05 (q, 2H, J = 17 Hz), 3.82 (dd, 1H, J = 17 Hz), 4.05 (q, 2H, J = 17 Hz), 3.82 (dd, 1H, J = 17 Hz), 4.05 (q, 2H, J = 17 Hz), 3.82 (dd, 1H, J = 17 Hz), 4.05 (q, 2H, J = 17 Hz), 4.05 (q, 2

17.2 Hz and J = 5.2 Hz), 3.72 (dd, 1H, J = 17 Hz and J = 5.6 Hz), 3.40 (s, 1H), 3.08 (d, 1H, J = 17 Hz), 2.61 (dd, 1H, J = 18 Hz and J = 7.2 Hz), 1.61 (s, 9H), 1.54 (s, 9H), 1.05 (s, 9H), 0.26 (s, 6H).

To a solution of trifluoroacetic acid (33.3 ml) and H<sub>2</sub>O (2.7 ml) was added 2-(*tert*-butoxyoxalyl-amino)-5-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (0.70 g, 1.04 mmol). The solution was stirred at room temperature for 40 hours. The solvent was poured into ethyl ether (400 ml). The precipitate was filtered off and dried <u>in vacuo</u>, which afforded 450 mg (80 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>): δ 12.30 (s, 1H), 9.71 (s, 1H), 9.20 (s, 2H), 7.39 (t, 1H, J = 8.0 Hz), 6.99 (d, 1H, J = 8.0 Hz), 6.82 (d, 1H, J = 8.0 Hz), 4.52 (d, 1H, J = 16.8 Hz), 4.36 (d, 2H, J = 17 Hz), 4.22 (d, 2H, J = 17 Hz), 4.00 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 3.86 (s, 1H), 3.62 (d, 1H, J = 17 Hz), 2.81 (dd, 1H, J = 18 Hz and J = 7.2 Hz);

LC-MS:  $R_t = 1.20 \text{ min}$ ;  $m/z = 432 [M+H]^{+}$ 

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### **EXAMPLE 79**

5-(7-Benzyloxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid
To a solution of 2-(tert-butoxyoxalyl-amino)-5-(7-(tert-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (2.40 g, 3.34 mmol) in 10 % formic acid/methanol (50 ml) at room
temperature under nitrogen was added 10 % Pd/C (1.2 g). The mixture was stirred for 48 hours. The Pd/C was filtered off and the filtrate was evaporated in vacuo. The residue was dissolved in dichloromethane (10 ml) and the resulting solution was poured into hexane. The precipitate was

filtered off (1.3 g) and the filtrate was evaporated in vacuo. The residual foam (1.1 g) was taken into dichloromethane (50 ml) and treated with ditert-butyl-dicarbonate (1.1 g, 5.0 mmol) and saturated sodium bicarbonate (20 ml). The mixture was stirred for 2 hours and the organic layer was separated and dried (MgSO<sub>4</sub>). The solvent was evaporated in vacuo and the residue was chromatographed using a gradient of 10-30% ethyl acetate/Hexane as eluent, which afforded 175 mg of 2-(tert-butoxyoxalyl-amino)-5-(7-hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-carboxylic acid di-tert-butyl ester.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.55 (s, 1H), 8.53 (s, 1H), 7.37 (t, 1H, J = 7.6 Hz), 6.92 (d, 1H, J = 7.6 Hz), 6.83 (d, 1H, J = 7.6 Hz), 4.95 (s, 1H), 4.84 (d, 1H, J = 16.4 Hz), 4.72 (d, 1H, J = 16.0 Hz), 4.56 (d, 1H, J = 16.0 Hz), 4.28 (d, 1H, J = 17.6 Hz), 4.13 (m, 1H), 3.68 (s, 0.5H), 3.42 (s, 0.5H), 3.16-2.94 (m, 2H), 1.62 (s, 9H), 1.61 (s, 9H), 1.26 (s, 9H).

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To a solution of 2-(*tert*-butoxyoxalyl-amino)-5-(7-hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-carboxylic acid di-*tert*-butyl ester (16 mg, 0.025 mmol) in N,N-dimethylformamide (0.5 ml) under nitrogen was added sodium hydride (1.0 mg, 0.026 mmol) at room temperature. The solution was stirred for 2 hours and followed by addition of benzyl bromide (5.9 ml, 0.050 mmol). The solution was stirred for 16 hours, diluted with ethyl acetate (20 ml) and washed with 0.5 N hydrochloric acid solution (2 x 10 ml), saturated sodium bicarbonate (2 x 10 ml), brine (10 ml), dried (MgSO<sub>4</sub>), and filtered.

The solvent was removed in vacuo. The residue was chromatographed using a gradient of 10-20 % ethyl acetate/Hexane as eluent, which afforded 14 mg (76 %) of 5-(7-benzyloxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(tert-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-carboxylic acid di-tert-butyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.49 (s, 1H), 7.48 (d, 2H, J = 7.2 Hz), 7.35 (m, 3H), 7.28 (d; 1H, J = 7.2 Hz), 6.97 (d, 1H, J = 7.6 Hz), 6.80 (d, 1H, J = 7.6 Hz), 5.32 (s, 2H), 4.97 (m, 2H), 4.82-4.62 (m, 2H), 4.45-4.15 (m, 2H), 3.68 (s,

0.5H), 3.48 (s, 0.5H), 3.16-2.94 (m, 2H), 1.62 (s, 9H), 1.60 (s, 9H), 1.26 (s, 9H).

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (2.7 ml) was added 5-(7-benzyloxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(*tert*-butoxyoxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-carboxylic acid di-*tert*-butyl ester (14 mg, 0.019 mmol). The solution was stirred at room temperature for 40 hours. The reaction mixture was poured into ethyl ether (20 ml). The precipitate was filtered off and dried <u>in vacuo</u> affording 8.0 mg (68 %) of the <u>title compound</u> as a solid trifluoroacetate.  $^1$ H-NMR (DMSO-d<sub>6</sub>):  $\delta$  12.25 (s, 1H), 9.28 (s, 1H), 9.02 (s, 1H), 7.53 (m, 3H), 7.39 (t, 2H, J = 7.6 Hz), 7.13 (d, 1H, J = 7.6 Hz), 7.11 (d, 1H, J = 8.4 Hz), 5.27 (m, 2H), 4.54 (d, 1H, J = 17.2 Hz), 4.38 (d, 2H, J = 17.6 Hz), 4.22 (m, 2H), 4.00 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 3.86 (s, 1H), 3.64 (d, 1H, J = 17.2 Hz), 2.81 (dd, 1H, J = 18 Hz and J = 7.2 Hz); LC-MS: R<sub>t</sub> = 2.96 min; m/z: 522 [M+H]<sup>+</sup>

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5-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml) was added 2-(*tert*-butoxyoxalyl-amino)-5-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (11 mg, 0.014 mmol). The solution was stirred at room temperature for 16 hours.

The reaction mixture was poured into ethyl ether (20 ml). The precipitate

was filtered off and dried <u>in vacuo</u>, which afforded 7.0 mg (79 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>): δ 12.39 (s, 1H), 9.95 (s, 1H), 9.75 (s, 2H), 7.42 (t, 1H, J = 8.0 Hz), 7.30 (s, 2H), 7.02 (d, 1H, J = 7.2 Hz), 6.96 (s, 2H), 6.85 (d, 1H, J = 7.2 Hz), 4.95-3.65 (m, 11H), 3.76 (s, 3H). LC-MS: R<sub>t</sub> = 1.93 min, m/z: 553 [M+H]<sup>+</sup>

### **EXAMPLE 81**

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5-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a stirred solution of 2-amino-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (15 mg, 0.028 mmol) in tetrahydrofuran (1.0 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (27 mg, 0.11 mmol) in tetrahydrofuran (1.0 ml). The mixture was stirred at room temperature for 24 hours. The solvent was removed in vacuo. The residue was taken into ethyl acetate (20 ml). The solution was washed with 0.5 N hydrochloric acid solution (2 x 10 ml), saturated sodium bicarbonate (2 x 10 ml) and brine (10 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was removed in vacuo. The residue was chromatographed using a gradient of 10-25 % ethyl acetate/hexane as eluent, which afforded 17 mg (93 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.54 (s, 1H), 7.86 (m, 2H), 7.40 (m, 2H), 7.08 (d, 2H, J = 8.4 Hz), 6.72 (d, 2H, J = 8.4 Hz), 4.08 (dd, 1H, J = 13.6 Hz and J = 8.8 Hz), 3.94 (d, 1H, J = 16.8 Hz), 3.82 (d, 1H, J = 12.8 Hz), 3.78 (s, 3H), 3.92 (s, 3H), 3.70-3.56 (m, 3H), 3.53 (d, 1H, J = 12.8), 2.93 (dd, 1H, J = 16.8 Hz and J = 4.8 Hz), 2.75 (dd, 1H, J = 18.0 Hz and J = 5.6 Hz), 1.61 (s, 9H), 1.58 (s, 9H).

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml) was added 2-(*tert*-butoxyoxalyl-amino)-5-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (15 mg, 0.023 mmol). The solution was stirred at room temperature for 40 hours. The reaction mixture was poured into ethyl ether (20 ml). The precipitate was filtered off and dried <u>in vacuo</u>, which afforded 13 mg (87 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>): δ 12.38 (s, 1H), 7.89 (d, 4H, J = 11.2 Hz), 7.18 (s, 2H), 6.85 (s, 2H), 4.20-3.60 (m, 9H), 3.71 (s, 3H); LC-MS: R<sub>t</sub> = 2.05 min, m/z: 550 [M+H]<sup>†</sup>

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## EXAMPLE 82

7-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (80 mg, 0.20 mmol) and diisopropyl ethylamine (35 μl, 0.40 mmol) in acetonitrile (10 ml) at room temperature was added a solution of 6-bromomethyl-2-(*tert*-butyl-dimethyl-silanyloxy)-benzoic acid ethyl ester (69 mg, 0.20

mmol). The solution was stirred for 12 hours at room temperature and the solvent was evaporated in vacuo. The residue was dissolved in ethyl acetate (50 ml) and washed with water, 1 N hydrochloric acid, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was chromatographed on silica gel column eluted with 20 % ethyl acetate/hexane to yield 42 mg (33 %) of 2-amino-7-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  7.64 (d, 1H, J = 8.8 Hz), 7.39 (t, 1H, J = 8.0 Hz), 7.10-6.80 (m, 5H), 6.09 (s, 2H), 5.0-4.2 (m, 4H), 3.80 (s, 3H), 3.66-2.92 (m, 3H), 1.55 (s, 9H), 1.04 (s, 9H), 0.22 (s, 6H).

To a stirred solution of 2-amino-7-(7-(tert-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-15 thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (40 mg, 0.060 mmol) in tetrahydrofuran (1 ml) was added imidazol-1-yl-oxo-acetic acid tert-butyl ester (59 mg, 0.30 mmol) in tetrahydrofuran (1 ml). The mixture was stirred at room temperature for 24 hours. The solvent was removed in 20 vacuo. The residue was dissolved in ethyl acetate (20 ml) and the solution was washed with 0.5 N hydrochloric acid (2 x 20 ml), saturated sodium bicarbonate (2 x 20 ml), brine (20 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was removed in vacuo and the residue was chromatographed using a gradient of 10-20 % ethyl acetate/Hexane as eluent, which afforded 40 mg (83 %) of 2-(tert-butoxyoxalyl-amino)-7-(7-(tert-butyl-25 dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxybenzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.52 (s, 1H), 7.37 (t, 1H, J = 8.0 Hz), 6.97 (d, 2H, J = 8.4 Hz), 6.94 (d, 1H, J = 8.0 Hz), 6.83 (d, 1H, J = 8.0 Hz), 6.54 (d, 1H, J = 8.4 Hz), 4.26 (d, 1H, J = 16.8 Hz), 3.93-3.84 (m, 2H), 3.77 (d, 1H, J = 16.8 Hz), 3.69 (s, 3H), 3.66-3.48 (m, 3H), 3.42-3.32 (m, 1H), 2.95 (dd, 1H, J = 14.4 Hz and J = 4.8 Hz), 2.92-2.82 (m, 1H), 2.73 (dd, 1H, J = 14.4 Hz and

J = 4.8 Hz), 1.60 (s, 9H), 1.59 (s, 9H), 1.02 (s, 9H), 0.22 (d, 6H, J = 1.6 Hz).

To a solution of 2-(*tert*-butoxyoxalyl-amino)-7-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (4.0 mg, 5.1 μmol) in 10 % formic acid/methanol (1 ml) at room temperature under nitrogen was added 10 % Pd/C (4 mg). The mixture was stirred for 1 hour. The Pd/C was filtered off and the filtrate was evaporated <u>in vacuo</u> to afford 2.8 mg (82 %) of 2-(*tert*-butoxyoxalyl-amino)-7-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-5H-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.45 (s, 1H), 8.05 (s, 1H), 7.39 (t, 1H, J = 8.0 Hz), 6.99 (d, 1H, J = 8.0 Hz), 6.79 (d, 1H, J = 8.0 Hz), 4.50 (d, 1H, J = 17.2 Hz), 4.45 (d, 1H, J = 17.2 Hz), 4.24 (d, 1H, 8.4 Hz), 4.03 (dd, 1H, J = 16.0 Hz and J = 7.2 Hz), 3.78-3.68 (m, 2H), 3.38-3.28 (m, 1H), 3.21 (d, 1H, J = 18.8 Hz), 3.08-2.98 (m, 1H), 1.57 (s, 9H), 1.56 (s, 9H), 0.98 (s, 9H), 0.15 (d, 6H, J = 1 Hz).

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml) was added 2-(*tert*-butoxyoxalyl-amino)-7-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-5*H*-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (2.8 mg, 0.0042 mmol). The solution was stirred at room temperature for 16 hours. The solvent was removed <u>in vacuo</u> and the residue was washed with dichloromethane affording 1.8 mg (79 %) of the <u>title compound</u> as a solid trifluoroacetate.

1-NMR (DMSO-d<sub>6</sub>): δ 12.30 (s, 1H), 9.76 (s, 1H), 9.40 (s, 1H), 8.95 (s, 1H), 7.40 (t, 1H, J = 7.6 Hz), 7.00 (d, 1H, J = 7.6 Hz), 6.83 (d, 1H, J = 7.6 Hz), 4.92 (s, 1H), 4.54 (d, 1H, J = 18.4 Hz), 4.40 (d, 2H, J = 18.4 Hz), 4.08-4.00 (m, 1H), 3.91 (d, 1H, J = 15.2 Hz), 3.60 (s, 2H), 3.06 (s, 2H);

LC-MS: R<sub>1</sub>: 1.41 min, m/z: 432 [M+H]<sup>+</sup>

### **EXAMPLE 83**

7-(7-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml) was added 2-(*tert*-butoxyoxalyl-amino)-7-(7-(*tert*-butyl-dimethyl-silanyloxy)-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (10 mg, 0.013 mmol). The solution was stirred at room temperature for 16 hours. The solvent was removed <u>in vacuo</u> and the residue was washed with dichloromethane, which afforded 6.8 mg (92 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>): δ 12.35 (s, 1H), 9.90 (s, 1H), 9.70 (s, 2H), 7.41 (t, 1H, J = 8.0 Hz), 7.28 (s, 2H), 7.04 (d, 1H, J = 7.2 Hz), 6.92 (s, 2H), 6.83 (d, 1H, J = 7.2 Hz), 4.90-3.60 (m, 11H), 3.80 (s, 3H). LC-MS:  $R_t = 1.92 \text{ min, m/z: } 552 \text{ [M+H]}^+$ 

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# EXAMPLE 84 O OH ON OH OH

7-(1,3-Dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid
To a stirred solution of 2-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-

carboxylic acid *tert*-butyl ester (10 mg, 0.019 mmol) in tetrahydrofuran (1.0 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (18 mg, 0.092 mmol) in tetrahydrofuran (1.0 ml). The mixture was stirred at room temperature for 24 hours. The solvent was removed <u>in vacuo</u>. The residue was dissolved in ethyl acetate (20 ml) and washed with 0.5 N hydrochloric acid solution (2 x 10 ml), saturated sodium bicarbonate (2 x 10 ml), brine (10 ml), dried (MgSO<sub>4</sub>), and filtered. The solvent was removed <u>in vacuo</u> and the residue was chromatographed using a gradient of 10-25 % ethyl acetate/hexane as eluent, which afforded 11 mg (89 %) of 2-(*tert*-

butoxyoxalyl-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 12.54 (s, 1H), 7.76 (m, 4H), 6.82 (d, 2H, J = 11.6 Hz), 6.33 (d, 2H, J = 11.6 Hz), 4.02 (d, 1H, J = 14.4 Hz), 3.98 (d, 1H, J = 14.4 Hz), 3.62 (s, 3H), 3.62-3.54 (m, 2H), 3.48-3.34 (m, 2H), 3.02-2.70 (m, 3H), 1.60 (s, 9H), 1.59 (s, 9H).

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml) was added 2-(*tert*-butoxyoxalyl-amino-7-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (10 mg, 0.015 mmol). The solution was stirred at room temperature for 16 hours. The solvent was removed <u>in vacuo</u> and the residue was washed with dichloromethane, which afforded 6.8 mg (80 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>): δ 12.38 (s, 1H), 7.86 (m, 4H), 6.82 (s, 2H), 6.30 (s, 2H), 4.00-2.86 (m, 9H), 3.58 (s, 3H); LC-MS:  $R_t = 2.02 \text{ min}$ ; m/z: 550 [M+H]<sup>+</sup>

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7-(((5-Benzyloxy-1*H*-indole-2-carbonyl)amino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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2-Amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (0.50 g; 1.2 mmol) was dissolved in N,N-dimethylformamide (20 ml). 1-Hydroxy-7azabenzotriazole (0.19 g; 1.3 mmol) and N-(3-dimethylaminopropyl)-N'ethylcarbodiimide hydrochloride (0.26 g; 1.3 mmol) and diisopropylethylamine (0.23 ml; 1.3 mmol) were added and the mixture was stirred for 15 min. 5-Benzyloxyindole (0.36 g; 1.3 mmol) was dissolved in N,Ndimethylformamide (20 ml) and added. Diisopropylethylamine (0.23 ml; 1.3 mmol) was added and the mixture was stirred overnight. The solvent was removed in vacuo, the residue dissolved in dichloromethane (30 ml) and the organic phase washed with an aqueous solution of sodium hydrogencarbonate (15 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered and the solvent removed in vacuo. The residue was chromatographed on silica using ethyl acetate/heptane (1:1) as eluent affording 569 mg of 2-amino-7-(((5-benzyloxy-1H-indole-2carbonyl)amino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid *tert-*butyl ester as an oil.

The <u>title compound</u> was prepared in a similar way as described in Example 48 using the last two steps.

MS: m/z: 669.4 [M+H]<sup>†</sup>

Calculated for  $C_{35}H_{32}N_4O_8S$ ,  $2/3xC_2HF_3O_2$ ,  $4/3xH_2O$  ;

C, 56.77%; H, 4.63%; N, 7.29%. Found:

C, 56.43%; H, 4.57%; N, 7.13%.

### **EXAMPLE 86**

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7-(((6-Bromo-2-p-tolyl-quinoline-4-carbonyl)amino)methyl)-6-(4-methoxybenzyl)-2-(oxalyl-amino)- 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acidThe <u>title compound</u> was prepared in a similar way as in Example 84 using 6-bromo-2-p-tolyl-quinoline-4-carboxylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material. LC-MS: m/z: 745.2 [M+H]<sup>+</sup>

Calculated for C<sub>36</sub>H<sub>31</sub>BrN<sub>4</sub>O<sub>7</sub>S, 2xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>;

C, 49.44%; H, 3.42%; N, 5.77%. Found:

15 C, 49.19%; H, 3.59%; N, 6.00%.

### EXAMPLE 87

6-(4-Methoxy-benzyl)-7-(((5-methyl-2-phenyl-2*H*-[1,2,3]triazole-4-carbonyl)amino)-methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-

20 c]pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as in Example 84 using 5-methyl-2-phenyl-2*H*-[1,2,3]triazole-4-carboxylic acid and 2-amino-7-

aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z: 605.2 [M+H]<sup>+</sup>

5 Calculated for  $C_{29}H_{28}N_6O_7S$ ,  $1.3xC_2HF_3O_2$ ,  $1.7xH_2O$ ;

C, 48.14%; H, 3.94%; N, 10.94%. Found:

C, 48.35%; H, 4.19%; N, 10.68%.

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### **EXAMPLE 88**

7-(((1*H*-Indole-3-carbonyl)amino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

- The <u>title compound</u> was prepared in a similar way as in Example 84 using 3-indole-carboxylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.
- 20 LC-MS: m/z: 563.2 [M+H]<sup>+</sup>
  Calculated for C<sub>28</sub>H<sub>26</sub>N<sub>4</sub>O<sub>7</sub>S, 5/3xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>;
  C, 49.63%; H, 3.82%; N, 7.35%. Found:
  C, 50.00%; H, 3.71%; N, 7.44%.

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7-((4-Ethoxy-2-hydroxy-benzoylamino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as in Example 84 using 4-ethoxy-2-hydroxy-benzoic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z: 584 [M+H]<sup>+</sup>

10 HPLC: (B6): 23.8 min.

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### **EXAMPLE 90**

7-((4-Benzoylamino-benzoylamino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

The title compound was prepared in a similar way as in Example 84 using 4-benzoylaminobenzoic acid and 2-amino-7-aminomethyl-6-(4-methoxy-

benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z: 643.1 [M+H]<sup>+</sup>

Calculated for  $C_{33}H_{30}N_4O_8S$ ,  $3xC_2HF_3O_2$ ;

5 C, 47.57%; H, 3.38%; N, 5.69%. Found:

C, 47.34%; H, 3.55%; N, 5.62%.

# **EXAMPLE 91**

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7-(((Biphenyl-4-carbonyl)amino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as in Example 84 using 4-phenylbenzoic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z: 599.0 [M+H]<sup>+</sup>

20 Calculated for  $C_{32}H_{29}N_3O_7S$ ,  $2xC_2HF_3O_2$ ,  $1xH_2O$ ;

C, 51.13%; H, 3.93%; N, 4.97%. Found:

C, 52.02%; H, 4.02%; N, 5.16%.

7-(((1*H*-Indole-2-carbonyl)amino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as in Example 84 using indole-2-carboxylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z: 563.2 [M+H]<sup>+</sup>

10 HPLC (B6)  $R_t = 23.07 \text{ min.}$ 

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# **EXAMPLE 93**

7-((3-Biphenyl-4-yl-acryloylamino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c)pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as in Example 84 using 3-biphenyl-4-yl-acrylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

LC-MS: m/z:  $626.2 [M+H]^{+}$ HPLC (B6) R<sub>t</sub> = 28.74 min.

Calculated for C<sub>34</sub>H<sub>31</sub>N<sub>3</sub>O<sub>7</sub>S, 2xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>;
 C, 53.46%; H, 3.90%; N, 4.92%. Found:
 C, 53.89%; H, 4.23%; N, 5.08%.

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### **EXAMPLE 94**

6-(4-Methoxy-benzyl)-7-(((5-methoxy-1*H*-indole-2-carbonyl)amino)-methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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The <u>title compound</u> was prepared in a similar way as in Example 84 using 5-methoxyindole-2-carboxylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

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LC-MS: m/z:  $593.2 [M+H]^{+}$ HPLC (B6) R<sub>t</sub> = 21.81 min.

### EXAMPLE 95

7-((4-Benzyl-benzoylamino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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The <u>title compound</u> was prepared in a similar way as in Example 84 using 4-benzylbenzoic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

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LC-MS: m/z: 614.2 [M+H] $^{+}$ HPLC (B6) R<sub>t</sub> = 27.23 min. Calculated for C<sub>33</sub>H<sub>31</sub>N<sub>3</sub>O<sub>7</sub>S, 1.5xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>, 1xH<sub>2</sub>O; C, 53.87%; H, 4.33%; N, 5.23%. Found:

15 C, 53.92%; H, 4.24%; N, 5.18%.

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### **EXAMPLE 95**

6-(4-Methoxy-benzyl)-7-(((naphthalene-1-carbonyl)amino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid The <u>title compound</u> was prepared in a similar way as in Example 84 using 1-napthylcarboxylic acid and 2-amino-7-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as the starting material.

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LC-MS: m/z: 574.0  $[M+H]^+$ HPLC (B6)  $R_t = 22.51$  min.

Calculated for C<sub>30</sub>H<sub>27</sub>N<sub>3</sub>O<sub>7</sub>S, 2xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>; 10 C, 50.94%; H, 3.65%; N, 5.24%. Found: C, 51.39%; H, 3.79%; N, 5.16%.

### **EXAMPLE 96**

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6-(4-Methoxy-benzyl)-5-((2-naphthalen-2-yl-ethylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

A solution of 2-naphthalen-2-yl-ethanol (1.02 g, 5.8 mmol), 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) (9 mg, 0.058 mmol) and sodium bromide (0.65 g, 6.4 mmol) in a mixture of toluene (18 mL), ethyl acetate (18 mL), and water (3mL) was cooled to 0 °C and added dropwise over 1 hour a solution containing the following: sodium hypochlorite (17.2 mL, 0.37 M, 6.4 mmol) and sodium hydrogencarbonate (1.46 g, 17.4 mmol). The reaction mixture was stirred at 0 °C for 10 min., and the phases separated. The aqueous layer was extracted with ethyl acetate (150 mL). The combined organic phases were washed with a solution of potassium

iodone (0.2 g) in 10 % aqueous potassium hydrogensulfate (150 mL),

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water (150 mL), brine (150 mL), dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo to provide 980 mg of a 3:1 mixture of naphthalen-2-yl-acetaldehyde and 2-naphthalen-2-yl-ethanol.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 9.81 (t, 1H, J = 1.5 Hz), 7.92-7.80 (m, 3H), 7.68 (bs, 1H), 7.55-7.42 (m, 3H), 3.87 (d, 2H, J = 1.5 Hz).

To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (290 mg, 0.71 mmol) in 1,2-dichloroethane (3 ml) was added the above mixture of 2-naphthyl-acetaldehyde (100 mg, 0.59 mmol), sodium triacetoxyborohydride (190 mg, 0.88 mmol) and the mixture was stirred at room temperature under nitrogen for 2.5 hours. The crude reaction mixture was quenched with saturated sodium bicarbonate (50 ml) and the solution extracted with ethyl acetate (100 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and concentrated <u>in vacuo</u> providing a foam, which was taken directly to the next step. LC-MS showed that 2-amino-6-(4-methoxy-benzyl)-5-((2-naphthalen-2-yl-ethylamino)-methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was the major component.

20 LC-MS: m/z: 558.1 [M+H] $^{\dagger}$ , R<sub>f</sub> = 2.23 min.

To a solution of 2-amino-6-(4-methoxy-benzyl)-5-((2-naphthalen-2-ylethylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester in tetrahydrofuran (3 ml) was added di-*tert*-butyldicarbonate (188 mg, 0.85 mmol) and *N,N*-dimethylformamide (18 mg, 0.14 mmol). The reaction was stirred at room temperature for 7 hours under nitrogen. The crude reaction mixture was diluted with dichloromethane (50 ml) and washed with water (50 ml) and brine (50 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo affording a foam, which was used without further purification in the next step.

LC-MS showed that 2-amino-5-((*tert*-butoxycarbonyl-(2-naphthalen-2-yl-ethyl)-amino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was the major component. R<sub>f</sub>= 2.74, m/z: 658.1 [M+H]<sup>+</sup>, Calculated: 657.4.

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To crude 2-amino-5-((tert-butoxycarbonyl-(2-naphthalen-2-yl-ethyl)amino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid tert-butyl ester was added dichloromethane (5 ml) and imidazol-1-yl-oxo-acetic acid tert-butyl ester (400 mg, 1.78 mmol) and the reaction mixture stirred at room temperature for 12 hours. The crude reaction mixture was added to dichloromethane (50 ml) and washed with water (50 ml) and brine (50 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo. The residue was purified by flash chromatography using a mixture of dichloromethane/ethyl acetate (10.1) as eluent, which afforded 20.3 mg (39 % over tree steps) of 2-(tertbutoxyoxalyl-amino)-5-((tert-butoxycarbonyl-(2-naphthalen-2-yl-ethyl)amino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid *tert*-butyl ester as a foam. <sup>1</sup>H NMR (CDCl<sub>3</sub>) δ 7.99-7.92 (m, 3H), 7.88 (s, 1H), 7.68-7.57 (m, 3H), 7.45 (d, 2H, J = 7.8 Hz), 6.99 (d, 2H, J = 8.1 Hz), 3.90-3.75 (m, 7H), 3.56-3.42(m, 5H), 3.19-3.13 (m, 2H), 2.88-2.82 (m, 2H), 1.79 (s, 9H), 1.71 (s, 18H); LC-MS: m/z: 786.2 [M+H] $^{+}$ , R<sub>f</sub> = 3.03 min.

To a solution of 2-(*tert*-butoxyoxalyl-amino)-5-((*tert*-butoxycarbonyl-(2-naphthalen-2-yl-ethyl)-amino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (20 mg, 0.03 mmol) in dry dichloromethane (200  $\mu$ l) at 0 °C was added 50 % trifluoroacetic acid in dichloromethane (2.5 ml). The reaction was stirred for 14 hours at room temperature and then concentrated <u>in vacuo</u>. The resultant solid was re-suspended in dichloromethane, filtered, and dried <u>in vacuo</u> to provide 13 mg (90 %) of the <u>title compound</u> as a solid.

1 H NMR-(DMSO-d<sub>6</sub>)- $\delta$  9-15 (s, 1H), 8:09=8:01 (m, 3H), 7:93 (s, 1H), 7:68-7.57 (m, 3H), 7:45 (d, 2H, J = 7.8 Hz), 6.99 (d, 2H, J = 8.1 Hz), 4.18-4.12

(m, 2H), 3.90-3.75 (m, 7H), 3.56-3.42 (m, 3H), 3.19-3.13 (m, 2H), 2.88-2.82 (m, 2H);

LC-MS: m/z: 574.7 [M+H]<sup>+</sup>, R<sub>f</sub> = 1.36 min.

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### **EXAMPLE 97**

5-((2-Benzo[1,3]dioxol-5-yl-acetylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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To a mixture of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (300 mg, 0.74 mmol), benzo[1,3]dioxol-5-yl-acetic acid (134 mg, 0.74 mmol), 1-hydroxybenzotriazole hydrate (111 mg, 0.82 mmol), and *N*,*N*-diisopropylethylamine (258  $\mu$ L, 1.48 mmol) in acetonitrile (5 ml) at room temperature was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (157 mg, 0.82 mmol). The reaction mixture was stirred for 16 hours and the solvent evaporated <u>in vacuo</u>. The residue was taken into ethylacetate (50 ml), washed with water, 1 N hydrochloric acid, saturated sodium bicarbonate, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was subjected to flash chromatography using a gradient of 10-20% ethylacetate/hexanes as eluent, which afforded 268 mg (64 %) of 2-amino-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 6.95 (bs, 2H), 6.75-6.85 (m, 5H), 5.96 (bs, 2H), 5.95 (s, 2H), 3.81 (s, 3H), 3.75-3.30 (m, 5H), 3.53 (s, 2H), 3.18 (bs, 2H), 2.82 (d, 1H, J = 17 Hz), 2.52 (d, 1H, J = 17 Hz).

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To a solution of 2-amino-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (133 mg, 0.235 mmol) in tetrahydrofuran (1 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (100 mg, 0.51 mmol).

- The mixture was stirred at room temperature for 24 hours. The solvent was removed in vacuo. The residue was taken into ethyl acetate (50 ml) washed with saturated sodium bicarbonate, brine, dried (Na<sub>2</sub>SO<sub>4</sub>) and filtered. The solvent was removed in vacuo and the residue was chromatographed using a gradient of 10-20% ethyl
- acetate/dichloromethane, which afforded 130 mg (80 %) of 2-(*tert*-butoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.
  - <sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 12.50 (s, 1H), 7.95-7.75 (m, 7H), 5.96 (s, 2H), 3.81 (s, 3H), 3.80-3.40 (m, 5H), 3.15 (bs, 2H), 2.90 (d, 1H, J = 17 Hz), 2.58 (d, 1H, J = 17 Hz), 1.61 (s, 9H), 1.60 (s, 9H).

A solution of 2-(tert-butoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-ylacetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid tert-butyl ester (130 mg, 0.188 mmol) in tetrahydrofuran (2 ml) was passed through a Raney Ni bed (120 mg, 50% Raney Ni-water washed with methanol (6 ml) and tetrahydrofuran (6 ml) and dried before use). The Raney Ni bed was washed with tetrahydrofuran (10 ml). The filtrate and washes were combined and the solvent evaporated in vacuo. The residue was dissolved in 10% formic acid/methanol (6 ml) and stirred with 10% Pd/C (120 mg) for 13 hours. Saturated sodium bicarbonate solution (60 ml) was added to the solution. The mixture was extracted with dichloromethane. The extracts were combined, dried (Na<sub>2</sub>SO<sub>4</sub>) and filtered. The solvent was removed in vacuo and the residue was washed with 50% hexane/diethyl ether to afford 62 mg (57 %) of 2-(tert-butoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-ylacetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 12.59 (s, 1H), 6.80-6.70 (m, 3H), 5.96 (s, 2H), 4.05 (q, 2H, J = 15 Hz), 3.85-3.60 (m, 2H), 3.25-3.00 (m, 4H), 2.58 (m, 1H), 1.61 (s, 9H), 1.59 (s, 9H);

LC-MS:  $R_t = 1.75 \text{ min, m/z}$ : 574 [M+H]<sup>+</sup>.

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A solution of 2-(*tert*-butoxyoxalyl-amino)-5-((2-benzo[1,3]dioxol-5-yl-acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (62 mg, 0.11 mmol) in 50% trifluoroacetic acid-dichloromethane (2 ml) was left in an open flask over the weekend and then the solvent was removed <u>in vacuo</u>. The residue was washed with dichloromethane and the solid filtered off affording 39 mg (62 %) of the title compounds as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.39 (s, 1H), 9.18 (bs, 1H), 9.10 (bs, 1H), 8.35 (s, 1H), 6.83 (d, 1H, J = 1.2 Hz), 6.82 (d, 1H, J = 8.4 Hz), 6.70 (dd, 1H, J = 1.5 Hz), 5.96 (s, 2H), 4.38 (d, 1H, J = 14 Hz), 4.28 (m, 1H), 3.60-3.40 (m, 4H), 3.16 (d, 2H, J = 14 Hz), 2.80 (dd, 1H, J = 14 Hz and J = 11 Hz);

LC-MS: R<sub>t</sub> = 1.11 min, m/z: 462 [M+H]<sup>+</sup>.

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### **EXAMPLE 98**

5-((2-Dibenzofuran-2-yl-ethyl)amino)methyl)-6-(4-methoxy-benzyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid To a solution of 2-dibenzofuran-2-yl-ethanol (200 mg, 0.94 mmol) and 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) (2 mg, 0.009 mmol) in dichloromethane (2 mL) was added an aqueous solution of sodium bromide (97 mg in 1.3 mL of water for a 0.7M solution, 0.94mmol) and cooled to 0 °C. To this mixture was added dropwise over 30 min., a

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solution containing the following: sodium hypochlorite (1.4 mL, 0.74 M, 1.03 mmol) and sodium hydrogencarbonate (120 mg, 1.4 mmol) and water (1.4 mL). The reaction mixture was stirred at 0 °C for 0.5 hour and allowed to warm to room temperature. The organic phase and aqueous layer were separated and the aqueous layer extracted with dichloromethane (20 mL). The combined organic phases were washed with a solution of potassium iodone (0.2 g) in 10% aq. Potassium hydrogensulfate (20 mL), water (20 mL), brine (20 mL), dried (MgSO<sub>4</sub>) filtered, and concentrated in vacuo to provide 198 mg of a 5:1 mixture of dibenzofuran-2-yl-acetaldehyde and 2-dibenzofuran-2-yl-ethanol as an oil. <sup>1</sup>H-NMR (CDCl<sub>3</sub>): δ 9.80 (t, 1H, *J* = 1.5 Hz), 8.02 (d, 2H, *J* = 8.2 Hz), 7.71 (bs, 1H), 7.75-7.42 (m, 4H), 3.82 (d, 2H, *J* = 1.5 Hz).

To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (340 mg, 0.85 mmol) in 1,2-dichloroethane (3 ml) was added the above mixture of dibenzofuran-2-yl-acetaldehyde (150 mg, 0.70 mmol), and sodium triacetoxyborohydride (225 mg, 1.07 mmol) and the mixture was stirred at room temperature under nitrogen for 2.5 hours. The crude reaction mixture was quenched with saturated sodium bicarbonate (50 ml) and the solution extracted with ethylacetate (100 ml). The organic phase dried (MgSO<sub>4</sub>), filtered, and the solvent evaporated <u>in vacuo</u>. The crude residue was taken directly to the next step. LC-MS showed that 2-amino-5-((2-dibenzofuran-2-yl-ethylamino)methyl]-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was the major component in the crude mixture: m/z: 598.1 [M+H]<sup>+</sup>, R<sub>f</sub> = 2.40 min).

Crude 2-amino-5-((2-dibenzofuran-2-yl-ethylamino)methyl]-6-(4-methoxybenzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was diluted in tetrahydrofuran (3 ml) and di-*tert*-butyl dicarbonate (262 mg, 1.20 mmol) and 4-(*N*,*N*-dimethylamino)pyridine (25 mg, 0.20 mmol) were added. The reaction was stirred at room temperature for 7 hours under nitrogen. The crude reaction mixture was added to

dichloromethane (50 ml) and washed with water (50 ml) and brine (50 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered, and concentrated <u>in vacuo</u>. The residue was used directly in the next step. LC-MS showed that 2-amino-5-((*tert*-butoxycarbonyl-(2-dibenzofuran-2-yl-ethyl)amino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was the major component in the crude: R<sub>f</sub> = 2.76, m/z: 698.2 [M+H]<sup>†</sup>.

To compound 2-amino-5-((tert-butoxycarbonyl-(2-dibenzofuran-2-ylethyl)amino)-methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-10 c]pyridine-3-carboxylic acid tert-butyl ester was added dichloromethane (5ml) and imidazol-1-yl-oxo-acetic acid tert-butyl ester (420 mg, 2.12 mmol). The reaction mixture was stirred at room temperature for 12 hours. The crude reaction mixture was added to dichloromethane (50 ml) and washed with water (50 ml) and brine (50 ml). The organic phase was dried 15 (MgSO<sub>4</sub>), filtered, and concentrated in vacuo. The residue was subjected to flash chromatography using a mixture of dichloromethane/ethyl acetate (10:1) as eluent, which afforded 35.2 mg (51 % over 3 steps) of 2-(tertbutoxyoxalyl-amino)-5-((tert-butoxycarbonyl-(2-dibenzofuran-2-ylethyl)amino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-20 c]pyridine-3-carboxylic acid *tert*-butyl ester as a foam. <sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.95-7.90 (m, 3H), 7.84 (s, 1H), 7.68-7.57 (m, 3H), 7.45 (d, 2H, J = 7.8 Hz), 6.95 (m, 3H), 3.90-3.75 (m, 7H), 3.56-3.42 (m, 5H),3.19-3.13 (m, 2H), 2.88-2.82 (m, 2H), 1.79 (s, 9H), 1.71 (s, 18H); LC-MS:  $R_f = 3.03 \text{ min, m/z}$ : 826.2 [M+H]<sup>+</sup>. 25

To a solution of 2-(*tert*-butoxyoxalyl-amino)-5-((*tert*-butoxycarbonyl-(2-dibenzofuran-2-yl-ethyl)amino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (28 mg, 0.034 mmol) in dry dichloromethane (200 μL) at 0 °C was added 50% trifluoroacetic acid in dichloromethane (2.5 ml). The reaction was stirred for 14 hours at room-temperature and then concentrated in vacuo. The resultant solid was re-suspended in dichloromethane, filtered, and dried in

vacuo, which afforded 22 mg (90 %) of the title compound as a solid trifluoroacetate.

 $^{1}\text{H-NMR}$  (DMSO-d<sub>6</sub>)  $\delta$  9.15 (s, 1H), 8.11-8.21 (m, 3H), 7.93 (s, 1H), 7.68-7.57 (m, 3H), 7.45 (d, 2H, J = 7.8 Hz), 6.99 (d, 2H, J = 8.1 Hz), 4.18-4.12 (m, 2H), 3.90-3.75 (m, 7H), 3.56-3.42 (m, 3H), 3.19-3.13 (m, 2H), 2.88-2.82 (m, 2H);

LC-MS:  $R_f = 3.03$ , m/z:  $614.7 [M+H]^+$ .

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# EXAMPLE 99

6-(4-Methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1H-indol-3-yl)acetylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3clpyridine-3-carboxylic acid

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To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (202 mg, 0.50 mmol), in N,N-dimethylformamide (4 ml) was added 5-methoxy-2methyl-3-indole acetic acid (170 mg, 0.74 mmol), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide, hydrochloride (150 mg, 0.75 mmol), and 1hydroxybenzotriazole (105 mg, 0.74 mmol). The mixture was stirred at room temperature for 12 hours. The crude reaction mixture was diluted with dichloromethane (100 ml) and washed with water (100 ml), brine (100 ml), dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo, which afforded 2amino-6-(4-methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1H-indol-3yl)acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-25 carboxylic acid tert-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.16 (d, 2H, J = 10.8 Hz), 6.99 (d, 1H, J = 2.5 Hz), 6.94 (m, 1H), 6.85 (dd, 1H, J = 8.4 Hz and J = 1.2 Hz), 6.78 (dd, 1H, J = 8.3 Hz and J = 1.2 Hz), 6.65 (m, 3H), 6.57 (m, 4H), 3.57 (t, 4H, J = 3.0 Hz), 3.53 (m, 6H), 3.59-3.29 (m, 5H), 3.12-2.92 (m, 4H), 2.39 (s, 3H), 1.6 (s, 9H); LC-MS  $R_t = 2.19$ , m/z: 605 [M+H]<sup> $\dagger$ </sup>.

To a solution of 2-amino-6-(4-methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1*H*-indol-3-yl)acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (96 mg, 0.5 mmol) in dichloromethane (5 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (583 mg, 3.0 mmol) and the reaction stirred at room temperature for 24 hours. The mixture was then concentrated <u>in vacuo</u>. The residue was purified by flash column chromatography (25% ethylacetate/dichloromethane) to give 53 mg (15 %) of 2-(*tert*-butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1*H*-indol-3-yl)acetylamino)methyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 7.16 (d, 2H, J = 10.8 Hz), 6.99 (d, 1H, J = 2.5 Hz), 6.94 (m, 1H), 6.85 (dd, 1H, J = 8.4 Hz and J = 1.2 Hz), 6.78 (dd, 1H, J = 8.3 Hz and J = 1.2 Hz), 6.65 (m, 3H), 6.56 (m, 3H), 3.57 (m, 3H), 3.53 (m, 6H), 3.59-3.29 (m, 5H), 3.12-2.92 (m, 4H), 2.39 (s, 3H), 1.6 (s, 18H); LC-MS R<sub>t</sub> = 2.36 min, m/z: 733 [M+H]<sup>+</sup>.

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2-(*tert*-Butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-((2-(5-methoxy-2-methyl-1*H*-indol-3-yl)acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was dissolved in 50% trifluoroacetic acid/dichloromethane (3 ml) and stirred at room temperature for 48 hours. The solvent was removed <u>in vacuo</u> and the residual trifluoroacetic acid was removed under reduced pressure to give 17 mg (49 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 10.62 (s, 1H), 7.31 (s, 1H), 7.08 (d, 1H, J = 10.2 Hz), 6.93 (s, 2H), 6.58 (dd, 1H, J<sub>1</sub> = 5.25 Hz and J<sub>2</sub> = 2.8 Hz), 3.84-3.44 (m, 19H, partially obscured by solvent), 2.95 (s, 1H), 2.28 (s, 3H), 1.31 (s, 1H), 1.19 (s, 2H);

LC-MS-R<sub>t</sub> = 1.89-min, m/z: 621 [M+H]<sup>+</sup>.

### **EXAMPLE 100**

5-((2-(1*H*-Indol-3-yl)-2-oxo-acetylamino)methyl)-2-(Oxalyl-amino)-4,5,6,7-

5 <u>tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid</u>

To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (209 mg, 0.51 mmol) in dry *N*,*N*-dimethylformamide (4 ml) was added 3-indole-glyoxylic acid (141 mg, 0.74 mmol), 1-(3-dimethylaminopropyl)-3-

ethylcarbodiimide, hydrochloride (152 mg, 0.76 mmol), and 1-hydroxybenzotriazole (100 mg, 0.74 mmol). The mixture was stirred at room temperature for 16 hours, diluted with dichloromethane (100 ml) and washed with water (100 ml), brine (100 ml), dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo. The residue was subjected to flash

chromatography using a mixture of ethyl acetate/hexanes (2:5) as eluent, which afforded 143 mg (40 %) of 2-amino-5-((2-(1*H*-indol-3-yl)-2-oxo-acetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

LC-MS R<sub>t</sub> = 2.31 min, m/z: 574.9 [M+H]<sup>+</sup>.

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To a solution of 2-amino-5-((2-(1*H*-indol-3-yl)-2-oxo-acetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (143 mg, 0.25 mmol) in dichloromethane (5 ml) was added imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (144 mg, 0.75 mmol) and the flask was purged with nitrogen. After 24 hours an additional portion of imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (169 mg, 0.86 mmol) was added and the reaction mixture allowed stirred for an additional 24 hours. The mixture was then concentrated <u>in vacuo</u>. The residue was purified by flash chromatography using a mixture of ethyl

acetate/hexanes (2:5) as eluent, which afforded 101 mg (58 %) of 2-(tert-butyoxyoxalyl-amino)-5-((2-(1*H*-indol-3-yl)-2-oxo-acetylamino)methyl)-6-(4-

methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 9.23 (s, 1H), 9.07 (d, 1H, J = 3.6 Hz), 8.50 (d, 1H, J = 7.6 Hz), 8.15 (d, 1H, J = 4.0 Hz), 7.47 (d, 2H, J = 7.2 Hz), 7.38-7.27 (m, 6H), 6.89 (d, 2H, J = 8.8 Hz), 3.87-3.59 (m, 6H), 3.04 (dd, 2H, J = 23.6 Hz), 2.74 (dd, 2H, J = 22.4 Hz), 1.62 (s, 18H); LC-MS R<sub>t</sub> = 2.49 min, m/z: 703 [M+H]<sup>+</sup>.

2-(tert-Butyoxyoxalyl-amino)-5-((2-(1H-indol-3-yl)-2-oxoacetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-10 c]pyridine-3-carboxylic acid tert-butyl ester (101 mg, 0.143 mmol) was dissolved in dry tetrahydrofuran (6 ml) and passed through a pipette, plugged with cotton containing Raney 2800 Nickel (0.38 g). The pipette was flushed with dry tetrahydrofuran (6 ml) and the filtrate was concentrated in vacuo. Pd on carbon (10%, 102 mg, source: Avocado) 15 and formic acid (10% in methanol, 5 ml) were added to the flask containing 2-(tert-Butyoxyoxalyl-amino)-5-((2-(1H-indol-3-yl)-2-oxoacetylamino)methyl)-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3clpyridine-3-carboxylic acid tert-butyl ester. After stirring for 18 hours, the solution was filtered through a pad of celite and concentrated in vacuo. 20 The residue was diluted in ethyl acetate, washed with saturated sodium bicarbonate (2 x 25 ml), brine (2 x 25 ml), dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo. The residue was subjected to flash chromatography using a mixture of 10% methanol/dichloromethane as 25 eluent, which afforded 2-(tert-butyoxyoxalyl-amino)-5-((2-(1H-indol-3-yl)-2oxo-acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3carboxylic acid tert-butyl ester. <sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  9.23 (s, 1H), 9.07 (d, 1H, J = 3.6 Hz), 8.50 (d, 1H, J = 7.6 Hz), 8.15 (d. 1H, J = 4.0 Hz), 7.27 (s. 2H), 7.09 (d. 1H, J = 8.8 Hz), 6.81 (d, 1H, J = 8.8 Hz), 3.79 (s, 1H), 2.29 (s, 1H), 1.62-1.57 (m, 18H), 30 0.08 (s, 5H);

LC-MS:  $R_t = 2.17 \text{ min, m/z:} 583 \text{ [M+H]}^{+}$ .

The above 2-(*tert*-butyoxyoxalyl-amino)-5-((2-(1*H*-indol-3-yl)-2-oxo-acetylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester was dissolved in 50% trifluoroacetic acid/dichloromethane (3 ml) and stirred at room temperature for 18 hours. The solvent was removed <u>in vacuo</u> and residual trifluoroacetic acid was removed under reduced pressure affording 17.1 mg of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.28 (s, 2H), 9.26 (s, 1H), 9.13 (s, 1H), 8.83 (d, 1H, J = 2.8 Hz), 8.26 (d, 1H, J = 8.8 Hz), 7.55 (d, 1H, J = 4.8 Hz), 7.27 (d, 2H, J = 7.6 Hz), 4.42 (d, 1H, J = 15.2 Hz), 4.29 (d, 1H, J = 16.4 Hz), 3.76-3.22 (m, 4H, partially obscured by solvent), 2.91-2.834 (m, 1H), 1.23 (s, 1H); LC-MS: R<sub>t</sub> = 0.99 min, m/z 471.4 [M+H]<sup>+</sup>.

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# GENERAL CHIRAL SYNTHESIS 4-Oxo-1-((S)-1-phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester

Dichloromethane (1L) and mol sieves 3 Å (113 g) and amine (S)-(-)α-methyl-benzylamin (71,7 ml) were mixed in a 2 l three-necked bottle cooled to -5 °C (using a ethanol/water/ice bath). A 50 % solution of ethylglyoxylate in toluene (117,6 ml) was added drop wise over 20 min., keeping the temperature between -5 °C and 0 °C The mixture was stirred for 0.5 hour before it was cooled to -30 °C. Trifluoroacetic acid (45,2 ml) was added over 3-4 minutes. Boron trifluoride diethyl ether (69,8 ml) was added drop wise over 5 min at -55 °C. The ice bath was removed and the mixture was allowed to warm up to -45 °C whereupon 2-(trimethylsilyloxy)-1,3-butadiene (100 ml) was added drop wise over 10 minutes. During the addition the mixture was cooled and the temperature

kept below -20 °C. The above additions are all exothermic hence the cooling bath should have sufficient capacity to remove the heat generated during the rapid addition. The reaction mixture was stirred for 2 hours at -15 °C and 1 hour at 0 °C and then poured on ice/water and stirred for 15 minutes. Solid sodium hydrogen carbonate was added until pH 7-8. The mixture was stirred overnight at room temperature. The layers wee separated and the aqueous phase extracted with dichloromethane. The combined organic phases were filtered through a plug of silica eluting with dichloromethane. The relevant fractions were concentrated in vacuo. The residue was dissolved in hot heptane and cooled. This leaves a yellowish gummy material on the side of the flask and crystals starts forming. The heptane solution was heated again to dissolve crystals, leaving the gummy material on the side of the flask and the mixture was filtered hot. The heptane solution was cooled to room temperature and the precipitate was filtered off and dried in vacuo, which afforded 38 g of 4-oxo-1-((S)-1phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester as a solid.

The filtrate was put in a refrigerator and a second crop was formed which was less pure and needed recrystallization from heptane to yield another 7,5 g of 4-oxo-1-((S)-1-phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester.

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4,4-Diethoxy-1-((S)-1-phenyl-ethyl)-piperidine-(S)-2-carboxylic acid ethyl ester

The mother liquor from the above crystallization was concentrated in vacuo. 5.0 g of the resulting material (18.16 mmol) was dissolved in

ethanol (100 ml) and triethylorthoformate (26.9 g, 181.6 mmol) and paratoluensulphonic acid (6.9 g, 36.32 mmol) was added. The reaction was stirred at room temperature for 16 hours before the mixture was poured on aqueous sodium hydrogen carbonate (200 ml) and extracted with ethyl acetate (4 x 75 ml). The combined extracts were concentrated in vacuo and purified by column chromatography (SiO<sub>2</sub>, Flash 40, petrol ether-ethyl acetate 10:1). Collection of the first band ( $R_f = 0.68$ ) gave 1.14 g (18 %) of 4,4-diethoxy-1-((S)-1-phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester and collection of the second band ( $R_f = 0.4$ ) gave 3.60 g (57 %) of the title compound.

# 4,4-Diethoxy-1-((S)-1-phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester

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4-Oxo-1-((*S*)-1-phenyl-ethyl)-piperidine-(*R*)-2-carboxylic acid ethyl ester (11.0 g, 0.040 mmol) was dissolved in a 1:1 mixture of triethyl orthoformate and ethanol (140 ml) and *para*-toluene-4-sulphonic acid (15.2 g, 80 mmol) was added and the reaction mixture was stirred for 16 hours. The reaction mixture was neutralized with sodium bicarbonate (to pH 7-8), and extracted with dichloromethane (3 x 100 ml), dried (MgSO<sub>4</sub>), filtered and concentrated <u>in vacuo</u>. The residue was purified by column chromatography (SiO<sub>2</sub>, petrol ether/ethyl acetate 10:1), which afforded 12.0 g (86 %) of the <u>title compound</u> as an oil.

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4,4-Diethoxy-1-((S)1-phenyl-ethyl)-(R)-2-hydroxymethyl-piperidine

To a solution of 4,4-diethoxy-1-((*S*)-1-phenyl-ethyl)-piperidine-(*R*)-2-carboxylic acid ethyl ester (36.0 g, 0.103 mol) in dry diethyl ether (150 ml) was added a suspension of lithium aluminum hydride (5.88 g, 0.155 mol)

in dry diethyl ether (300 ml) under an atmosphere of nitrogen at such a rate that the solution gently reflux. The reaction mixture was stirred over night before it was cooled to 0 °C and ethyl acetate (30 ml) was added drop wise to destroy excess lithium aluminum hydride. After stirring for another 0.5 hour, water (12 ml) was added drop wise. After stirring for 10-15 min the precipitate was filtered off through celite and the filter cage was washed with plenty of diethyl ether. The filtrate was washed with brine (100 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo, which afforded 30 g (95 %) of the title compound as an oil.

15 4,4-Diethoxy-1-((S)-1-phenyl-ethyl)-(R)-2-phthalimidomethyl-piperidine

A solution of 4,4-Diethoxy-1-((*S*)1-phenyl-ethyl)-(*R*)-2-hydroxymethyl-piperidine (65.35 g, 0.213 mmol), triphenylphosphine (61.3 g, 0.234 mol) and phthalimide (34.4 g, 0.234 mol) in tetrahydrofuran (700 ml) cooled to 0 °C was added diethyl azodicarboxylate over the course of 1.5 hour. The reaction mixture was stirred at 0 °C for another 2 hours before the solvent was removed in vacuo. The residue was dissolved in hot heptane-toluene (3:2) (650 ml) before it was cooled on an ice bath. The precipitate consisting of triphenyl phosphine oxide was filtered off and washed with heptane. The filtrate was concentrated in vacuo and the residue subjected

to column chromatography using a mixture of toluene-ethyl acetate-heptane (3:1:3) as eluent. The solvent was evaporated <u>in vacuo</u> whereupon a viscous oil was obtained. Upon addition of light petrol ether the product crystallized to give 67.4 g (73 %) of the <u>title compound</u> as a solid.

## EXAMPLE 101

5-(R)-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

A mixture of 4,4-diethoxy-1-((*S*)-1-phenyl-ethyl)-(*R*)-2-phthalimidomethyl-piperidine (5.25 g, 12.0 mmol) and hydrazine hydrate (2.92 ml, 60 mmol) was stirred overnight in ethanol (100 ml) at room temperature. The solvent was removed <u>in vacuo</u> and the solid residue was extracted with refluxing diethyl ether. The diethyl ether fractions were combined and evaporated <u>in vacuo</u>, which afforded 3.94 g (94 %) of 4,4-diethoxy-1-((*S*)-1-phenyl-ethyl)-(*R*)-2-aminomethyl-piperidine as an oil.

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4,4-Diethoxy-1-((*S*)-1-phenyl-ethyl)-(*R*)-2-aminomethyl-piperidine (2.25 g, 7.37 mmol), and triethyl amine (1.49 g, 14.7 mmol) in acetonitrile (50 ml) was heated to 60 °C before 2-chlormethyl-6-methoxy-benzoic acid methyl ester (1.58 g, 7.37 mmol) in acetonitrile (25 ml) was added over the course of 1.5 hour. After addition the reaction mixture was stirred overnight at 60 °C. The solvent was removed <u>in vacuo</u> and the residue was dissolved in dichloromethane (50 ml) and washed with saturated sodium bicarbonate. After drying (MgSO<sub>4</sub>), filtration and evaporation of the solvent <u>in vacuo</u> the residue was subjected to flash column

chromatography (SiO<sub>2</sub>, ethyl acetate-light petrol ether (1:1)) to give 2.3 g

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(69 %) of 2-(*R*)-(7-methoxy-2,3-dihydro-isoindol-1-one-2-ylmethyl)-4,4-diethoxy-1-(1-(*S*)-phenyl-ethyl)-piperidine.

2-(R)-(7-Methoxy-2,3-dihydro-isoindol-1-one-2-ylmethyl)-4,4-diethoxy-1(1-(S)-phenyl-ethyl)-piperidine (2.0 g, 4.4 mmol) was dissolved in a ice cold mixture of trifluoroacetic acid and water (10 ml, 9:1) and stirred or 0.5 hour on an ice bath. The reaction mixture was poured on aqueous sodium carbonate (100 ml) and extracted with dichloromethane (2 x 50 ml). The organic phase was dried (MgSO<sub>4</sub>), filtered and evaporated in vacuo,
affording 1.67 g (100 %) of 2-(R)-(7-methoxy-2,3-dihydro-isoindol-1-one-2-ylmethyl)-4-oxo-1(1-(S)-phenyl-ethyl)-piperidine.

2-(*R*)-(7-Methoxy-2,3-dihydro-isoindol-1-one-2-ylmethyl)-4-oxo-1(1-(*S*)-phenyl-ethyl)-piperidine (1.67 g, 4.41 mmol), sulphur (0.155 g, 4.85 mmol), *tert*-butylcyanoacetate (0.684 g, 4.85 mmol), *N*-methylmorpholine (0.892 g, 8.82 mmol) and molecular sieves (4Å, 2 g) was heated to 50 °C in ethanol under an atmosphere of nitrogen for 16 hours. The reaction mixture was filtered through a plug (1 cm) of SiO<sub>2</sub>, the silica was washed with dichloromethane-ethyl acetate and the solvent was removed <u>in</u> vacuo. The resulting residue was subjected to column chromatography (Flash 40, SiO<sub>2</sub>, toluene-ethyl acetate (3:1)), which yielded 1.17 g (50 %) of 2-amino-5-(*R*)-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-*c*]pyridine-3-carboxylic acid *tert*-butyl ester and 2-amino-7-(*S*)-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-*c*]pyridine-3-carboxylic acid *tert*-butyl ester as a 3:1 mixture.

The above mixture of 5- and 7-regioisomers (1.17 g, 2.19 mmol) and imidazol-2-yl-oxo-acetic acid *tert*-butyl ester (1.29 g, 7.57 mmol) and triethylamine (0.66 g, 6.57 mmol) was stirred under an atmosphere of nitrogen in dichloromethane (25 ml) for 16 hours. The solvent was removed in vacuo and the residue was subjected to column chromatography (SiO<sub>2</sub>, Flash 40, ethyl acetate-petrol ether (1:1)).

Collection of relevant fractions gave 0.61 g (42 %) of 2-(*tert*-butoxyoxalyl-amin)-5-(*R*)-(7-methoxy-1-oxo-1,3-dihydro-isoindo-2-ylmethyl)-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester.

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2-(*tert*-Butoxyoxalyl-amin)-5-(*R*)-(7-methoxy-1-oxo-1,3-dihydro-isoindo-2-ylmethyl)-6-(1-(*S*)-phenyl-ethyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (0.60 g, 0.91 mmol) was stirred for 16 hours in a mixture of methanol and formic acid (10:1) (20 ml) in the presence of 10 % palladium on carbon (50 % water). The reaction mixture was filtered through a plug of Celite and washed with methanol. The volatiles were removed <u>in vacuo</u> and the residue was dissolved in dichloromethane (50 ml), washed with semi saturated aqueous sodium carbonate (50 ml), dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was purified by column chromatography (SiO<sub>2</sub>, Flash 40, ethyl acetate-methanol (100:15)), which afforded 0.36 g (71 %) of 2-(*tert*-butoxyoxalyl-amin)-5-(*R*)-(7-methoxy-1-oxo-1,3-dihydro-isoindo-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester.

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2-(*tert*-Butoxyoxalyl-amin)-5-(*R*)-(7-methoxy-1-oxo-1,3-dihydro-isoindo-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (349 mg, 0.63 mmol) was stirred for 16 hours in a mixture of trifluoroacetic acid and dichloromethane (1:1) (10 ml) whereupon diethyl ether (20 ml) was added. The precipitate was filtered off and washed with diethyl ether, which afforded 215 mg (61 %) of the <u>title compound</u> as a solid trifluoroacetate.

LC-MS:  $R_t = 1.17 \text{ min, m/z. } 446 \text{ [M+H]}^{+}$ Calculated for  $C_{20}H_{19}N_3O_7S$ ,  $C_2HF_3O_2$ ,  $0.5xH_2O_1$ 

30 C, 46.48%; H, 3.72%; N, 7.39%; Found: C, 46.45%; H, 3.97%; N, 7.43%;

#### **EXAMPLE 102**

5-(S)-(7-Methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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A solution of 4,4-diethoxy-1-((*S*)-1-phenyl-ethyl)-piperidine-(*S*)-2-carboxylic acid ethyl ester (35.98 g, 0.103 mol) in diethyl ether (150 ml) was added drop wise to a suspension of lithium aluminum hydride (5.88 g, 0.155 mol) in diethyl ether (300 ml) over the course of 1 hour. The reaction mixture was stirred at room temperature overnight before it was cooled on an ice bath and the reaction was quenched by dropwise addition of ethyl acetate (30 ml), followed by drop wise addition of water (12 ml) whereupon a gray precipitate was formed. The mixture was filtered through a plug of Celite and the filter cage was washed with plenty of diethyl ether. The filtrate was dried (MgSO<sub>4</sub>) before it was filtered and the solvent removed <u>in vacuo</u>, which afforded 24.5 g (79 %) of 4,4-diethoxy-1-(1-(*S*)-phenyl-ethyl)-(*S*)-2-hydroxymethyl-piperidine as an oil.

A suspension of 4,4-diethoxy-1-(1-(*S*)-phenyl-ethyl)-(*S*)-2-hydroxymethylpiperidine (20 g, 65 mmol), triphenylphosphine (18.76 g, 72 mmol) and phthalimide (10.52 g, 72 mmol) in tetrahydrofurane (200 ml) cooled to 0 °C was added diethyl azodicarboxylate (11.34 ml, 72 mmol) over the course of 1 hour. The reaction mixture was stirred at 0 °C for another 2 hours before the volatiles were removed in vacuo. The residue was dissolve in hot heptane-toluene (3:2) (100 ml) before it was cooled on an ice bath. The precipitate was filtered off and washed with heptane. The filtrate was concentrated in vacuo and the residue subjected to column chromatography using a mixture of toluene/ethyl acetate/heptane (3:1:3) as eluent. The solvent was evaporated in vacuo and the residue was crystallized by addition of light petrol ether (250 ml). The precipitate was

filtered off, which afforded 24 g (85 %) of 4,4-diethoxy-1-(1-(S)-phenylethyl)-2-(S)-phthalimidomethyl-piperidine as a solid.

4,4-Diethoxy-1-(1-(S)-phenyl-ethyl)-2-(S)-phthalimidomethyl-piperidine (4.0 g. 9.2 mmol) was dissolved in a mixture of trifluoroacetic acid and water (9:1) (100 ml) at 0 °C and stirred for 2 hours at this temperature. The mixture was basified with half saturated aqueous sodium carbonate, extracted with ethyl acetate and dried (MgSO<sub>4</sub>) for 2 hours. The solvent was removed in vacuo and the residue was dried in a vacuum own at 40 °C for to days. This afforded 3.23 g (98 %) of 4-oxo-1-(1-(S)-phenyl-ethyl)-2-(S)-phthalimidomethyl-piperidine pure without further purification (98 %). A mixture of 4-oxo-1-(1-(S)-phenyl-ethyl)-2-(S)-phthalimidomethylpiperidine (17.28 g, 47.73 mmol), tert-butylcyanoacetat (7.41 g, 52.17 mmol), sulphur (1.71 g, 52.17 mmol) and morpholine (8.31 g, 95.46 mmol) in ethanol (150 ml) was heated under an atmosphere of nitrogen at 50 °C. The volatiles were removed in vacuo and the residue was subjected to column chromatography on silica gel (heptane-ethyl acetate 5:1). The fractions consisting of a mixture of 5- and 7-isomer were collected and the solvent evaporated in vacuo. The residue was purified on a reverse phase (C<sub>18</sub>) column using a Flash 40 system. The residue was applied in a minimum volume of acetonitrile and eluted with 40 % acetonitrile in water containing 0.1 % trifluoroacetic acid. When the 5-isomer was collected the eluent was changed to 50 % acetonitrile in water with 0.1 % trifluoroacetic acid and the 7-isomer was collected. Yield of 2-amino-5-(S)-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester was 7.96 g and yield of 2-amino-7-(R)-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(S)phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester was 3.72 g (47 % total).

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2-Amino-5-(S)-(1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butylester (7.96 g, 15.4 mmol) and hydrazine hydrate (3.85 g, 77.0 mmol) in

ethanol (250 ml) was stirred for 16 hours at room temperature. The solvent was removed in vacuo and the solid residue was extracted with diethyl ether (3 x 200 ml). The fractions were combined and the solvent removed in vacuo to give 5.9 g (100 %) of 2-amino-5-(S)-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester.

2-Amino-5-(*S*)-aminomethyl-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-*c*]pyridine-3-carboxylic acid *tert*-butyl ester (0.55 g, 1.42 mmol) and triethylamine (396 μl, 2.84 mmol) was heated in acetonitrile (15 ml) under an atmosphere of nitrogen to 60 °C whereupon a solution of 2-chloromethyl-6-methoxy-benzoic acid methyl ester (0.32 g, 1.49 mmol) in acetonitrile (5 ml) was added dropwise over the course of 3 hours, keeping the reaction mixture at 60 °C. The reaction was allowed to cool to room temperature and was left for 16 hours before the solvent was evaporated <u>in vacuo</u>. The product was purified by column chromatography (SiO<sub>2</sub>, Flash 40, ethyl acetate-petrol ether) to give 400 mg (53 %) of 2-amino-5-(*S*)-(7-methoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-((*S*)-1-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-*c*]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

The <u>title compound</u> was obtained as a trifluoroacetate in a similar way as described in example 101 using the last three steps.

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### EXAMPLE 103

5-(S)-(4-Hydroxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic-acid

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3-Hydroxy-2-methylbenzoic acid (0.5 g, 3.2 mmol) was dissolved in HPLC grade methanol (5 ml) and cooled to 0 °C under nitrogen. Acetyl chloride (5 ml) was added dropwise. Once the addition was complete, the ice bath was removed and the reaction mixture allowed warming to room temperature over a period of 18 hours. The reaction was complete by tlc (R<sub>f</sub>=0.5, 1:1 ethyl acetate/hexanes) and quenched with saturated sodium bicarbonate. The reaction mixture was concentrated, diluted with dichloromethane and water and the layers separated. The aqueous layer was extracted with dichloromethane (3x). The organic layers were combined, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo, which afforded 0.5 g (91 %) of 3-hydroxy-2-methylbenzoic acid methyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.39 (dd, 1H, J = 8.1 Hz and J = 1.5 Hz), 7.09 (t, 1H, J = 8.1 Hz), 6.92 (dd, 1H, J = 8.1 Hz and J = 1.2 Hz), 5.11 (bs, 1H), 3.87 (s, 3H), 2.43 (s, 3H).

3-Hydroxy-2-methylbenzoic acid methyl ester (0.5 g, 3.01 mmol) in dichloromethane (15 ml) and *N*,*N*-diisopropylethylamine (1.57 ml, 9.03 mmol) was cooled to 0 °C under nitrogen. Chloromethyl methyl ether (0.46 ml, 6.02 mmol) was added dropwise and the reaction allowed warming to room temperature over a period of 18 hours. The reaction was judged to be 50 % complete by tlc (1.2 ethyl acetate/hexanes, I<sub>2</sub>) and therefore, *N*,*N*-diisopropylethylamine (1.57 ml, 9.03 mmol) was added, the reaction mixture cooled to 0 °C and chloromethyl methyl ether (0.46 ml, 6.02 mmol) added once more. The reaction mixture was warmed to room temperature and stirred for 5 hours. The reaction was quenched with water and the layers separated. The aqueous layer was extracted once with dichloromethane and the organic layers combined, dried (MgSO<sub>4</sub>), filtered, and concentrated in vacuo. The crude residue was purified by column chromatography (20 % ethyl acetate/hexanes) affording 0.44 g (69 %) of 3-methoxymethoxy-2-methyl-benzoic acid methyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.46 (dd, 1H, J = 7.6 Hz and J = 1.2 Hz), 7.21 (dd, 1H, J = 8 Hz and J = 1.2 Hz), 7.18 (d, 1H, J = 8 Hz), 5.21 (s, 2H), 3.88 (s, 3H), 3.48 (s, 3H), 2.46 (s, 3H).

To a mixture of 3-methoxymethoxy-2-methyl-benzoic acid methyl ester (0.44 g, 2.09 mmol) in carbon tetrachloride (10 ml) was added *N*-bromosuccinimide (0.39 g, 2.19 mmol) and 1,1'-azo bis(cyclohexane-carbonitrile) (0.051 g, 0.21 mmol). The mixture was heated at reflux for 3 hours, at which time the reaction was judged complete by tlc (1:4 ethyl acetate/hexanes). The reaction mixture was cooled to room temperature and concentrated in <u>vacuo to</u> a solid. The solid was recrystallized from hexane leaving 0.44 g (82 %) of 2-bromomethyl-3-methoxymethoxy-benzoic acid methyl ester as a solid.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.58 (dd, 1H, J = 6.8 Hz and J = 2.4 Hz), 7.33-7.29 (m, 2H), 5.30 (s, 2H), 5.07 (s, 2H), 3.94 (s, 3H), 3.52 (s, 3H).

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To a stirred mixture of 2-amino-5-(S)-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (0.24 g, 0.67 mmol) in acetonitrile (30 ml) was added N,Ndiisopropylethylamine (0.16 ml, 0.93 mmol) under nitrogen. 2-Bromomethyl-3-methoxymethoxy-benzoic acid methyl ester (0.16 g, 0.55 mmol) dissolved in acetonitrile, was added via syringe pump at a rate of 0.3 ml/hour. Once the addition was complete, the reaction mixture was stirred at room temperature for 24 hours. TIc analysis (1:1 ethyl acetate/hexanes) indicated the reaction to be complete. The volatiles were removed in vacuo and the resultant oil dissolved in ethyl acetate/water. The layers were separated and the aqueous layer extracted with ethyl acetate (3x). The organic layers were combined, dried (MgSO<sub>4</sub>), filtered and the solvebt evaporated in vacuo, which afforded 0.34 g (100 %) of 2-amino-5-(S)-(4methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(S)-phenylethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester, which was used without further purification in the next step. <sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  7.51 (d, 1H, J = 6.8-Hz), 7.42 (t, 2H, J = 7.6 Hz), 7.23= 7.17 (m, 5H), 5.93 (s, 2H), 5.25 (s, 2H), 4.23 (s, 2H), 4.12 (q, 1H, J = 7.2

Hz), 3.94 (m, 1H), 3.85 (q, 1H, J = 6.4 Hz), 3.66 (d, 1H, J = 16.4 Hz), 3.50 (s, 3H), 3.48-3.46 (m, 1H), 3.20 (dd, 1H, J = 14 Hz and J = 6 Hz), 2.94-2.87 (m, 1H), 2.60 (m, 1H), 1.49 (s, 9H), 1.36 (d, 3H, J = 6.4 Hz); LC-MS: m/z:  $564.1 \, [M+H]^{+}$ .

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To a solution of 2-amino-5-(S)-(4-methoxymethoxy-1-oxo-1,3-dihydroisoindol-2-ylmethyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid tert-butyl ester (0.34 g, 0.60 mmol) in dichloromethane (10 ml) was added imidazol-1-yl-oxo-acetic acid tert-butyl ester (0.35 g, 1.8 mmol). The reaction mixture was stirred at room temperature for 18 hours and the solvent concentrated in vacuo. The residue was dissolved in ethyl acetate and washed with water (2 x 20 ml) and brine (2 x 25 ml). The organic layer was dried (MgSO<sub>4</sub>), filtered and the solvent evaporated in vacuo. The residue was subjected to flash chromatography using a mixture of ethyl acetate/hexanes (1:1) as eluent. The obtained residue was then subjected to chromatotron purification (1% methanol/ dichloromethane) and later to another flash chromatography (20 % ethyl acetate/hexanes to 25 % ethyl acetate/hexanes) to obtain 210 mg (50 %) of 2-(tert-butoxyoxalyl-amino)-5-(S)-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil.  $^{1}$ H-NMR (CDCl<sub>3</sub>)  $\delta$  12.50 (s, 1H), 7.51 (dd, 1H, J = 6.8 Hz and J = 1.2 Hz), 7.42 (t, 2H, J = 8 Hz), 7.25-7.17 (m, 5H), 5.23 (s, 2H), 4.24 (q, 2H, J = 16.8 Hz), 4.08 (d, 1H, J = 16.8 Hz), 4.01 (dd, 1H, J = 14 Hz and J = 8.8Hz), 3.89 (d, 1H, J = 17.6 Hz), 3.82 (q, 1H, J = 6.8 Hz), 3.56 (q, 1H, J =6.4 Hz), 3.51 (s, 3H), 2.28 (dd, 1H, J = 14 Hz and J = 6.4), 2.98-2.92 (m, 1H), 2.69 (d, 1H, J = 17.2), 1.56 (s, 9H), 1.54 (s, 9H), 1.38 (d, 3H, J = 6.8Hz); LC-MS: m/z: 692.5 [M+H]<sup>+</sup>.

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To a solution of 2-(*tert*-butoxyoxalyl-amino)-5-(*S*)-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (0.16 g,

0.23 mmol) in formic acid (10 % in methanol, 5 ml total) was added 10% palladium on carbon (85 mg, source: Avacado) and the reaction mixture allowed to stir at room temperature. After 6 hours, tlc (1:1 ethyl acetate/hexanes) analysis indicated reaction complete. The reaction mixture was filtered through a pad of celite and concentrated in vacuo. The crude product was purified via flash chromatography (gradient: 3% isopropyl alcohol/dichloromethane to 5 % isopropyl alcohol) to provide 0.11 g (82 %) of 2-(*tert*-butoxyoxalyl-amino)-5-(S)-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-

methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.
 <sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ 12.50 (bs, 1H), 7.48 (dd, 1H, J = 7.6 Hz and J = 0.8 Hz), 7.38 (t, 1H, J = 8 Hz), 7.22 (dd, 1H, J = 8 Hz and J = 0.8 Hz), 5.24 (s, 2H), 4.50 (q, 2H, J = 17.3 Hz), 4.02-3.90 (m, 2H), 3.74 (ddd, 2H, J = 34 Hz, J = 13.6 Hz and J = 5.6 Hz), 3.49 (s, 3H), 3.24 (m, 1H), 2.97 (ddd, 1H,

5 Hz, J = 13.6 Hz and J = 5.6 Hz), 3.49 (s, 3H), 3.24 (m, 1H), 2.97 (ddd, 1H) J = 20 Hz, J = 4.4 Hz and J = 2.8 Hz), 2.50 (m, 1H), 1.59 (s, 9H), 1.51 (s, 9H);

LC-MS; m/z: 587.8 [M+H]<sup>+</sup>.

2-(tert-Butoxyoxalyl-amino)-5-(S)-(4-methoxymethoxy-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (0.11 g, 0.18 mmol) was dissolved in neat trifluoroacetic acid (4 ml) and stirred at room temperature for 48 hours. The reaction mixture was concentrated in vacuo and the resultant solid washed with dichloromethane several times affording 100 mg (83 %) of the title compound as a solid trifluoroaceatet.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.29 (bs, 1H), 10.13 (s, 1H), 9.29 (bs, 1H), 9.10 (bs, 1H), 7.32 (t, 1H, J = 7.6 Hz), 7.17 (d, 1H, J = 7.2 Hz), 7.01 (d, 1H, J = 8 Hz), 4.52 (d, 1H, J = 17.2 Hz), 4.40-4.22 (m, 3H), 4.05 (dd, 1H, J = 14.4 Hz and J = 0.6 Hz), 2.60 (dm, 1H), 2.60 (dm, 1H), 2.80 (dm, 1H), 2.80 (dm, 1H), 3.80 (dm, 1H)

30 Hz and J = 9.6 Hz), 3.90 (bs, 1H), 3.69 (dm, 1H), 3.22 (dm, 1H), 2.80 (dm, 1H);

LC-MS:-m/z:-432-2 [M+H]\*:---

#### **EXAMPLE 104**

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2-(S)-(Oxalyl-amino)-5-((4-phenoxy-benzylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

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A solution of 2-amino-5-(S)-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (500 mg, 1.29 mmol) and 4-phenoxybenzaldehyde (256 mg, 1.29 mmol) was heated to 50 °C in ethanol (50 ml) for 1 hour in the presence of molecular sieves (4 A, 5 ml). The reaction mixture was cooled on an ice bath before sodium borohydride (98 mg, 2.59 mmol) was added in three portions over 45 min. The cooling bath was removed and the reaction mixture was allowed to reach room temperature. The mixture was filtered through a plug of Celite and the filter cage was washed with dichloromethane (3 x 25 ml). The solvent was removed in vacuo and the residue was redissolved in ethyl acetate (50 ml), washed with sodium bicarbonate (50 ml) and dried (MgSO<sub>4</sub>). The solvent was removed in vacuo before the residue was redissolved in acetonitrile (20 ml). Triethylamine (130 mg, 1.29 mmol), ditert-butyl dicarbonate (282 mg, 1.29 mmol) and 4-(N,N-dimethylamino)pyridine (5 mg, cat.) was added and the reaction mixture was stirred for 16 hours at room temperature. The volatiles were removed in vacuo and ethyl acetate (50 ml) was added and the solution was washed with saturated sodium bicarbonate (50 ml) and dried (MgSO<sub>4</sub>). The crude product was purified by column chromatography (SiO2, petroleum etherethyl acetate (9:1)) to give 325 mg (38% overall) of 2-amino-5-(S)-((4phenoxy-benzylamino)methyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester.

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The <u>title compound</u> was obtained as a trifluoroacetate in a similar way as described in example 96 using the last three steps.

Oxalation: Standard procedure (16 hours, 82 %)

Hydrogenolysis: standard procedure (Pd/C, 10% Pd, methanol-formic

5 acid, 16 hours, ((10:1)) (82% yield)

TFA cleavage: Standard procedure. Yield 150 mg (87%).

LC-MS m/z:  $482 [M+H]^{+}$ ,  $R_t = 1.87 min$ Calculated for  $C_{24}H_{23}N_3O_6S$ ,  $2x(C_2HF_3O_2)$ 

10 C, 47.40%, H, 3.55%; N, 5.92%; Found:

C, 47.47%; H, 3.87%; N, 5.88%;

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#### **EXAMPLE 105**

5-(S)-((4-Acetylamino-benzylamino)-methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

- The <u>title compound</u> was prepared as a trifluoroacetate in a similar way as described in Example 96 using 2-amino-5-(S)-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester and *N*-(4-formyl-phenyl)acetamide as the starting material.
- Calculated for C<sub>20</sub>H<sub>22</sub>N<sub>4</sub>O<sub>6</sub>S, 1.5xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>, 1.5xH<sub>2</sub>O
   C, 43.78%; H, 3.99%; N, 8.88%; Found:
   C, 44.20%; H, 4.43%; N, 8.75%;

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#### **EXAMPLE 106**

5 7-(S)-((Acetyl-(4-phenoxy-benzyl)amino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

A solution of 2-amino-7-(S)-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic aicd tert-butyl ester (500 mg, 1.29 mmol) and 4-phenoxybenzaldehyde (256 mg, 1.29 mmol) was heated to 50 °C in ethanol (50 ml) for 1 hour in the presence of molecular sieves (4 A, 5 ml). The reaction mixture was cooled on an ice bath before sodium borohydride (98 mg, 2.59 mmol) was added in three portions over 45 min. The cooling bath was removed and the reaction mixture was allowed to reach room temperature. The mixture was filtered through a plug of Celite and the filter cage was washed with dichloromethane (3 x 25 ml). The solvent was removed in vacuo and the residue was redissolved in ethyl acetate (50 ml), washed with sodium bicarbonate (50 ml) and dried (MgSO<sub>4</sub>). The solvent was removed in vacuo before the product was dissolved in dichloromethane (10 ml). The solution was cooled on an ice bath before di-isopropyl-ethyl amine (101 mg, 1.29 mmol) was added followed by drop wise addition of acetyl chloride (101 mg, 1.29 mmol) in dichloromethane (1 ml). The reaction mixture was stirred 1 hour at 0 °C and the solution was washed with sodium bicarbonate (10 ml) and dried (MgSO<sub>4</sub>). The crude product was purified by flash column chromatography (SiO<sub>2</sub>, ethyl acetate-petrol ether 1:3) to give 320 mg (41%) of 7-(S)-((acetyl-(4-phenoxy-benzyl)amino)methyl)-2-amino-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine=3-carboxylic acid

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The <u>title compound</u> was obtained as a trifluoroacetate in a similar way as described in example 96 using the last three steps.

Oxalation: Standard procedure (Yield 69%)

Hydrogenolysis and trifluoroacetic acid cleavage in one step, Standard procedure (Overall yield 6%)

LC-MS m/z = 524 [M+H]<sup>+</sup>, R<sub>t</sub> = 2.58 min Calculated for  $C_{26}H_{25}N_3O_7S$ ,  $C_2HF_3O_2$ , 0.5xH<sub>2</sub>O C, 52.01%; H, 4.21%; N, 6.50%; Found: C, 51.82%; H, 4.34%; N, 6.36%.

#### **EXAMPLE 107**

15 <u>7-(S)-((Acetyl-benzyl-amino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid</u>

A solution of 2-amino-7-(*S*)-aminomethyl-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-*c*]pyridine-3-carboxylic aicd *tert*-butyl ester (400 mg, 1.03 mmol) and benzaldehyde (105 mg, 1.03 mmol) was heated to 50 °C in ethanol (20 ml) for 1 hour in the presence of molecular sieves (4 A, 7 ml). The reaction mixture was cooled on an ice bath before sodium borohydride (78 mg, 2.06 mmol) was added in three portions over 45 min. The cooling bath was removed and the reaction mixture was allowed to reach room temperature. The mixture was filtered through a plug of Celite and the filter cage was washed with dichloromethane (3 x 25 ml). The solvent was removed <u>in vacuo</u> and the residue was redissolved in ethyl acetate (50 ml), washed with sodium bicarbonate (50 ml) and dried (MgSO<sub>4</sub>). The solvent was removed <u>in vacuo</u> before the product was dissolved in dichloromethane (20 ml). The solution was cooled on an ice

bath before di-isopropyl-ethyl amine (267 mg, 2.06 mmol) was added followed by drop wise addition of acetyl chloride (81 mg, 1.03 mmol) in dichloromethane (1 ml). The reaction mixture was stirred 1 hour at 0 °C before sodium bicarbonate (20 ml) was added. The mixture was extracted with dichloromethane (2 x 10 ml) and the combined organic fractions were dried (MgSO<sub>4</sub>). The crude product was purified by flash column chromatography (petrol ether/ethyl acetate (3:1)), which afforded 250 mg (46 %) of 7-(S)-((acetyl-benzyl-amino)methyl)-2-amino-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester.

The <u>title compound</u> was obtained as a trifluoroacetate in a similar way as described in example 96 using the last three steps.

Oxalation: Standard procedure (54%)
Hydrogenolysis: Standard procedure (methanol-formic acid (10:1)) Yield
38 mg (26%)
Trifluoroacetic acid cleavage: Standard procedure 33 mg (80%)

LC-MS m/z: 432 [M+H]<sup>+</sup>, R<sub>t</sub> = 1.52 min
Calculated for C<sub>20</sub>H<sub>21</sub>N<sub>3</sub>O<sub>6</sub>S x 1.5xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>, 2xH<sub>2</sub>O
C, 43.26%; H, 4.18%; N, 6.58%; Found:
C, 43.19%; H, 3.86%; N, 6.46%.

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#### EXAMPLE 108

5-(S)-((1,1-Dioxo-1*H*-benzo[d]isothiazol-3-ylamino)methyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of (S)-2-amino-5-aminomethyl-6-(1-(S)-phenyl-ethyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (1.0 g, 2.58 mmol) in dichloromethane (10 ml) at 0 °C was added N, Ndiisopropylethylamine (0.54 ml, 5.16 mmol). A solution of 3-chlorobenzo[d]isothiazole 1,1-dioxide (0.52 g, 2.58 mmol) in dichloromethane 5 (10 ml) was then added dropwise and stirred for 30 min. The solution was warmed to room temperature and washed with water and dried (MgSO<sub>4</sub>). The solvent was then removed in vacuo. The residue was taken into dichloromethane (15 ml) and imidazol-1-yl-oxo-acetic acid tert-butyl ester (1.0 g, 5.16 mmol) was added. The solution was stirred for 2 hours. The 10 solvent was removed in vacuo. The residue was taken into ethyl acetate (100 ml). The solution was washed with 0.5 N hydrochloric acid solution, saturated sodium bicarbonate and brine, dried (MgSO<sub>4</sub>) and filtered. The solvent was removed in vacuo. The residue was chromatographed using a mixture of 0-5% ethyl acetate/dichloromethane as eluent, which afforded 15 0.6 g (34 %) of 2-(tert-butoxyoxalyl-amino)-5-(S)-((1,1-dioxo-1H-benzo[d] isothiazol-3-ylamino)methyl)-6-(1-(S)-phenyl-ethyl)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil <sup>1</sup>H-NMR (CDCl<sub>3</sub>)  $\delta$  12.50 (s, 1H), 7.94-7.92 (m, 1H), 7.79-7.71 (m, 2H), 20 7.59-7.50 (m, 1H), 7.38-7.27 (m, 4H), 6.86 (d, 1H, J=4 Hz), 4.14 (d, 1H, J=12 Hz), 3.95 (d, 1H, J=17 Hz), 3.88 (q, 1H, J=6 Hz), 3.70-3.62 (m, 1H), 3.47 (t, 1H, J=13 Hz), 3.34-3.24 (m, 1H), 3.06 (dd, 1H, J=17, 6 Hz), 2.53

25 A solution of 2-(*tert*-butoxyoxalyl-amino)-5-(*S*)-((1,1-dioxo-1*H*-benzo[d] isothiazol-3-ylamino)methyl)-6-(1-(*S*)-phenyl-ethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (252 mg, 0.37 mmol) in tetrahydrofuran (12 ml) was passed through Raney Ni (0.95 g, 50% Raney Ni-Water washed with methanol (6 ml) and tetrahydrofuran (10 ml) and dried before use). The solvent was removed in vacuo. The residue was dissolved in acetic acid (7 ml) and hydrogenated with 10% Pd/C (250 mg) at 50 psi for 15 hours. The mixture was filtered and the filtrate was-added to saturated sodium bicarbonate solution. The solution was then

(d, 1H, J=17 Hz), 1.62 (s, 9H), 1.61 (s, 9H), 1.44 (d, 3H, J=7 Hz).

extracted with ethylacetate (3 x 100 ml). The extracts were combined and dried (MgSO<sub>4</sub>). The solvent was removed in vacuo. The residue was washed with diethyl ether affording 156 mg (73 %) of 2-(tert-butoxyoxalylamino)-5-(S)-((1,1-dioxo-1H-benzo[d] isothiazol-3-ylamino)-methyl)-

5 4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil.

 $^{1}$ H-NMR (CDCl<sub>3</sub>)  $\delta$  12.59 (s, 1H), 7.94-7.90 (m, 1H), 7.70-7.66 (m, 3H), 7.51 (s, 1H), 4.11 (d, 1H, J=12 Hz), 4.08 (q, 2H, J=17 Hz), 3.40 (dd, 1H, J=12, 6 Hz), 3.26-3.18 (m, 1H), 3.18 (d, 1H, J=17 Hz), 2.55 (dd, 1H, J=12,

10 6 Hz), 1.62 (s, 18H). LC-MS:  $R_t = 3.58 \text{ min, m/z: } 577 \text{ [M+H]}^{+}$ .

A solution of 2-(*tert*-butoxyoxalyl-amino)-5-(*S*)-((1,1-dioxo-1*H*-benzo[d] isothiazol-3-ylamino)methyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (149 mg, 0.26 mmol) in 50 % trifluoroacetic acid/dichloromethane (1 ml) was left in an open flask for 60 hours. The volatiles were removed <u>in vacuo</u> and the residue was washed with dichloromethane to yield 80 mg (54 %) of the <u>title compound</u> as a solid trifluoroacetate.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>) δ 12.29 (s, 1H), 9.80 (s, 1H), 9.51 (bs, 2H), 8.19 (d, 1H, J=5 Hz), 8.02-8.00 (m, 1H), 7.89-7.84 (m, 2H), 4.46 (d, 1H, J=16 Hz), 4.30 (d, 1H, J=16 Hz), 3.96-3.80 (m, 3H), 3.30 (d, 1H, J=17 Hz), 2.93 (dd, 1H, J=18, 10 Hz);

LC-MS:  $R_t = 0.68 \text{ min, m/z: } 465 \text{ [M+H]}^{+}$ .

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#### EXAMPLE 109

5-(4-Benzyloxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

The <u>title compound</u> was prepared in a similar way as described in Example 52 as a trifluoroacetate.

<sup>1</sup>H-NMR (400 MHz, DMSO-d<sub>6</sub>) δ 12.31 (s, 1H), 9.25 (bs, 2H), 7.80 (t, 1H, J = 8 Hz), 7.59-7.32 (m, 7H), 5.37 (s, 2H), 4.42-4.21 (m, 2H), 3.95-3.70 (m, 3H), 3.4-3.2 (obscured by water, 1H), 2.83-2.75 (m, 1H) LC-MS: R<sub>t</sub> = 2.16 min, m/z: 536.1 [M+H]<sup>+</sup>

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### EXAMPLE 110

5-(6-Methoxy-4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of 2-amino-5-aminomethyl-6-(4-methoxy-benzyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (57.4 mg, 0.142 mmol) and diisopropyl ethylamine (49 μl, 0.28 mmol) in acetonitrile (20 ml) at room temperature was added 2-bromomethyl-5-methoxy-isophthalic acid dimethyl ester (3.00 g, 7.45 mmol). The solution was stirred for 16 hours and the solvent evaporated <u>in vacuo</u>. The residue was taken into ethyl acetate (50 ml) and washed with water (2 x 20 ml), 1 N hydrochloric acid (20 ml), brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was chromatographed on silica gel column using a mixture of ethyl acetate/hexane (1:1) as eluent, which afforded 62 mg (71 %) of 2-amino-6-(4-methoxy-benzyl)-5-(6-methoxy-4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-

tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as a solid.

<sup>1</sup>H-NMR δ (CDCI<sub>3</sub>): δ 7.75 (d, 1H, J = 2.4 Hz), 7.55 (d, 1H, J = 2.4 Hz), 7.11 (bs, 2H), 6.74 (d, 2H, J = 8.0 Hz), 5.97 (s, 2H), 4.71 (d, 1H, J = 18.4 Hz), 4.62 (d, 1H, J = 18.4 Hz), 4.09 (m, 1H), 3.93 (s, 3H), 3.92 (s, 3H), 3.80 (m, 1H), 3.76 (s, 3H), 3.66-3.40 (m, 5H), 2.80 (d, 1H, J = 17.2 Hz), 2.64 (d, 1H, J = 17.2 Hz), 1.52 (s, 9H).

To a stirred solution of 2-amino-6-(4-methoxy-benzyl)-5-(6-methoxy-4methoxy-carbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-10 tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester (60 mg, 0.10 mmol) in tetrahydrofuran (1.0 ml) was added imidazol-1-yl-oxo-acetic acid tert-butyl ester (60 mg, 0.30 mmol) in tetrahydrofuran (1.0 ml). The mixture was stirred at room temperature for 24 hours. The solvent was removed in vacuo. The residue was taken into ethyl acetate (20 ml) and 15 washed with 0.5 N hydrochloric acid (2 x 10 ml), saturated sodium bicarbonate (2 x 10 ml) and brine (10 ml), dried (MgSO<sub>4</sub>) and filtered. The solvent was removed in vacuo and residue was chromatographed using a gradient ethyl acetate/hexane (10-25 %) as eluent, which afforded 40 mg (58 %) of 2-(tert-butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(6-methoxy-20 4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H-NMR δ (CDCI<sub>3</sub>): δ 12.54 (s, 1H), 7.75 (d, 1H, J = 2.4 Hz), 7.55 (d, 1H, J = 2.4 Hz), 7.10 (d, 2H, J = 8.0 Hz), 6.74 (d, 2H, J = 8.0 Hz), 4.74 (d, 1H, J = 18.4 Hz), 4.62 (d, 1H, J = 18.4 Hz), 4.05-3.90 (m, 2H), 3.94 (s, 3H), 3.92 (s, 3H), 3.82-3.48 (m, 5H), 3.77 (s, 3H), 2.95 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 2.67 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 1.61 (s, 9H), 1.58 (s, 9H).

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To a solution of 2-(*tert*-butoxyoxalyl-amino)-6-(4-methoxy-benzyl)-5-(6-methoxy-4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester

(38 mg, 0.055 mmol) in 10 % formic acid/methanol (1.0 ml) at room temperature under nitrogen was added 10 % Pd/C (38 mg). The mixture was stirred for 16 hours and the Pd/C was filtered off and the filtrate evaporated in vacuo. The residue was taken into dichloromethane (1.0 ml) poured into hexane. The precipitate was filtered off, affording 28 mg (82 %) of 2-(tert-butoxyoxalyl-amino)-5-(6-methoxy-4-methoxycarbonyl-1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as a solid.

<sup>1</sup>H-NMR δ (CDCl<sub>3</sub>): δ 12.45 (s, 1H), 10.90 (s, 1H), 10.69 (s, H), 7.73 (s, 1H), 7.42 (s, 1H), 4.85 (bs, 2H), 4.65 (bs, 1H), 4.42 (bs, 2H), 3.99 (bs, 2H), 3.96 (s, 3H), 3.89 (s, 3H), 3.35 (bs, 1Hz), 3.21 (bs, 1H), 1.62 (s, 9H), 1.56 (s, 9H).

To a solution of trifluoroacetic acid (0.5 ml) and dichloromethane (0.5 ml)
was added 2-(*tert*-butoxyoxalyl-amino)-5-(6-methoxy-4-methoxycarbonyl1-oxo-1,3-dihydro-isoindol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3c]pyridine-3-carboxylic acid *tert*-butyl ester (14 mg, 0.023 mmol). The
solution was stirred at room temperature for 40 hours. The reaction
mixture was poured into diethyl ether (20 ml). The precipitate was filtered
off, which afforded 10 mg (75 %) of the <u>title compound</u> as a solid
trifluoroacetate.

<sup>1</sup>H-NMR δ (DMSO-d<sub>6</sub>): δ 12.28 (s, 1H), 9.32 (s, 1H), 9.10 (s, 1H), 7.65 (d, 1H, J = 2.4 Hz), 7.50 (d, 1H, J = 2.4 Hz), 4.82 (d, 1H, J = 17.2 Hz), 4.65 (d, 1H, J = 17.6 Hz), 4.40 (d, 1H, J = 17.6 Hz), 4.30 (m, 1H), 4.10 (dd, 1H, J = 17.2 Hz and J = 5.2 Hz), 3.95 (s, 1H), 3.89 (s, 6H), 3.85 (d, 1H, J = 17.2 Hz), 2.81 (dd, 1H, J = 18 Hz and J = 7.2 Hz). LC-MS:  $R_t = 1.30$  min; m/z: 504 [M+H]<sup>+</sup>

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2-(Oxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3-carboxylic acid and

5 <u>2-(Oxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3-carboxylic acid</u>

To a solution of 2-aminomethyl-4-(2-spiro[1,3]dioxolane)-piperidine (193 mg, 1.12 mmol) and diisopropyl ethylamine (0.46 ml, 2.55 mmol) in acetonitrile (10 ml) cooled to 0 °C was added 2-chlorosulfonyl-benzoic acid methyl ester (278 mg. 1.18 mmol). The solution was stirred at 25 °C for 24 hours. Solvent was removed in vacuo and the residue was chromatographed using a mixture of ethyl acetate/hexane (1:3) as eluent, which afforded 199 mg (51 %) of 2-(4-(2-spiro[1,3]dioxolane)piperidin-2-ylmethyl)-1,1-dioxo-1,2-dihydro-1H-benzo[d]isothiazol-3-one as a solid.  $^{1}$ H-NMR (CDCl<sub>3</sub>):  $\delta$  7.99-7.96 (m, 1H), 7.66-7.53 (m, 3H), 5.01 (s, 1H), 4.73 (dm, 1H, J = 14.4 Hz), 4.06-3.93 (m, 6H), 3.25 (dd, 1H, J = 12.6 Hz), 3.06 (td, 1H, J = 13.5 Hz and J = 3.6 Hz), 1.93 (dd, 1H, J = 14.1 Hz and J = 5.7 Hz), 1.87 (dd, 1H, J = 14.1 Hz and J = 3.0 Hz), 1.76 (dd, 1H, J = 13.5 Hz and J = 5.1 Hz). LC-MS:  $R_{t}$  = 1.78; m/z: 339 [M+H] $^{+}$ .

2-(4-(2-Spiro[1,3]dioxolane)piperidin-2-ylmethyl)-1,1-dioxo-1,2-dihydro-1H-benzo[d]isothiazol-3-one (199 mg, 0.588 mmol) was dissolved in 2 M hydrochloric acid (12 ml) and the solution was heated to 50 °C for 24 hours. The volatiles were removed in vacuo and the residue (341 mg) was treated without further purification with saturated sodium carbonate (12 ml), dichloromethane (8 ml)-and-di-t-butyl-dicarbonate (1.64 g, 7.5 mmol). The mixture was stirred at 35 °C for 3 days and extracted with

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dichloromethane (30 ml). The organic solution was washed with saturated sodium bicarbonate, brine, dried (MgSO<sub>4</sub>), filtered and the solvent evaporated <u>in vacuo</u>. The residue was chromatographed on silica gel column using a mixture of ethyl acetate/hexane (1:3) as eluent, which afforded 115 mg (50 %) of 4-oxo-2-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-piperidine-1-carboxylic acid *tert*-butyl ester as an oil.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  8.06 (dd, 1H, J = 6.0, 1.8 Hz), 7.95-7.80 (m, 3H), 5.02 (bs, 1H), 4.35 (bs, 1H), 3.91(dd, 1H, J = 15.0 Hz and J = 8.4 Hz), 3.78 (dd, 1H, J = 14.7 Hz and J = 5.7 Hz), 3.53 (t, 1H, J = 10.8 Hz), 2.74 (dd, 1H, J = 15.0 Hz and J = 7.5 Hz), 2.60-2.38 (m, 3H), 1.32 (s, 9H).

To a solution of 4-oxo-2-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2ylmethyl)-piperidine-1-carboxylic acid tert-butyl ester (115 mg, 0.292 mmol) in absolute ethanol (5 ml) was added t-butyl cyanoacetate (57 µl, 15 0.41 mmol), sulfur (13 mg, 0.41 mmol) and morpholine (55 µl, 0.63 mmol). The solution was stirred at 50 °C for 14 hours. The solvent was evaporated in vacuo and the residue was chromatographed on silica gel column using a mixture of ethyl acetate/hexane (1:4) as eluent, which afforded 100 mg (62 %) of 2-amino-5-(1,1,3-trioxo-1,3-dihydro-1H-20 benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6dicarboxylic acid di-tert-butyl ester and 2-amino-7-(1,1,3-trioxo-1,3dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester as a mixture. <sup>1</sup>H-NMR (CDCl<sub>3</sub>):  $\delta$  8.10-8.00 (m, 1H), 7.98-7.77 (m, 2.8H), 7.66-7.58 (m, 25 0.2H), 6.11 (s, 0.4H), 6.06 (s, 0.6H), 5.59 (m, 0.2H), 5.39 (t, 0.3H, J = 5.7

'H-NMR (CDCl<sub>3</sub>):  $\delta$  8.10-8.00 (m, 1H), 7.98-7.77 (m, 2.8H), 7.66-7.58 (m, 0.2H), 6.11 (s, 0.4H), 6.06 (s, 0.6H), 5.59 (m, 0.2H), 5.39 (t, 0.3H, J = 5.7 Hz) 5.23 (bs, 0.3H), 5.04 (bs, 0.4H), 4.77 (d, 0.4H, J = 14.4 Hz), 4.60 (d, 0.4H, J = 14.4 Hz), 4.45-4.18 (m, 1H), 4.02-3.82 (m, 1.5H), 3.64 (dd, 0.5H, J = 14.7 Hz and J = 5.2 Hz), 3.30-2.60 (m, 2H), 1.54 (s, 7H), 1.53 (s, 2H), 1.26 (s, 7H), 1.21 (s, 2H).

To a stirred solution of the above 2-amino-5-(1,1,3-trioxo-1,3-dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,6-

dicarboxylic acid di-tert-butyl ester and 2-amino-7-(1,1,3-trioxo-1,3dihydro-1H-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester mixture (100 mg, 0.18 mmol) in acetonitrile (7 ml) was added imidazol-1-yl-oxo-acetic acid tertbutyl ester (290 mg, 1.46 mmol) in acetonitrile (1 ml). The mixture was stirred at room temperature for 16 hours. The solvent was removed in vacuo and the residue was taken into ethyl acetate. The solution was washed with 0.5 N hydrochloric acid solution, saturated sodium bicarbonate, brine, dried MgSO<sub>4</sub>) and filtered. The solvent was removed in vacuo and the residue was chromatographed on silicagel using a mixture 10 of ethyl acetate/hexane (1:4) as eluent, which provided 98 mg (80 %) of a mixture of 2-(tert-butoxyoxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1Hbenzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6dicarboxylic acid di-tert-butyl ester and 2-(tert-butoxyoxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-15 5*H*-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester as a solid. <sup>1</sup>H-NMR (CDCl<sub>3</sub>):. $\delta$  12.60 (s, 0.3H), 12.54 (s, 0.7H), 8.12-8.06 (m, 1H), 7.98-7.80 (m, 2.8H), 7.66-7.58 (m, 0.2H), 5.83 (bs, 0.1H), 5.61 (t, 0.2H), 5.40-4.54 (m, 0.9H), 4.53-4.40 (m, 0.8H), 4.02-3.70 (m, 1.42H), 3.66 (dd, 0.58H, J = 14.7 Hz and J = 5.2 Hz), 3.30-2.99 (m, 3H), 1.68 (s, 6H), 1.62 20 (s, 6H), 1.60 (s, 6H), 1.31 (s, 4.5H), 1.25 (s, 4.5H);

To a solution of trifluoroacetic acid (4 ml) and dichloromethane (2 ml) was added the mixture of 2-(*tert*-butoxyoxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester and 2-(*tert*-butoxyoxalyl-amino)-7-(1,1,3-trioxo-1,3-dihydro-1*H*-benzo[d]isothiazol-2-ylmethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (78 mg, 0.12 mmol). The solution was stirred at room temperature for 24 hours. The solvent was then evaporated <u>in vacuo</u>, which afforded 50 mg (72 %) of the <u>title compounds</u> as a mixture of trifluoroacetates.

LC-MS:  $R_t = 4.45$ ; m/z: 678 [M+H]<sup> $\dagger$ </sup>.

<sup>1</sup>H-NMR (DMSO-d<sub>6</sub>):  $\delta$  12.32 (s, 1H), 9.75-9.20 (m, 2H), 8.40 (t, 1H, J = 6.0 Hz), 8.22-8.02 (m, 3H), 5.03 (bs, 0.5H), 4.52 (d, 1H), 4.38-4.10 (m, 2H), 3.88 (bs, 0.5H), 3.70-3.64 (m, 0.5H), 3.44-3.34 (m, 0.5H), 3.20-2.90 (m, 2H).

5 LC-MS:  $R_t = 1.28 \text{ min, m/z: } 466 \text{ [M+H]}^{\dagger}$ 

#### **EXAMPLE 112**

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# 7-(R)-Carbamoyl-2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of 2-(S)-4-oxo-piperidine-1,2-dicarboxylic acid 1-tert butyl ester (18.4 g, 75.6 mmol) and triethylamine (12.65 mL, 90.79 mmol) in 15 tetrahydrofuran (50 mL) cooled to -20°C was added isobutylchloroformate (11.81 mL, 90.79 mmol) and the mixture was stirred for 10 min at -20°C before a 25 % solution of ammonia in water (100 mL) was added. The temperature was kept at -20°C for 30 min before the cooling bath was 20 removed and the reaction mixture was allowed to reach room temperature and stirring was continued for another hour. The reaction mixture was extracted with ethyl acetate (6 x 50 mL) and the combined organic phases were dried (MgSO<sub>4</sub>). The solvent was removed in vacuo and the residue was purified by column chromatography (SiO<sub>2</sub>, Flash 40, ethyl acetate) to 25 give 8.51 g (46 %) of 2-(S)-carbamoyl-4-oxo-piperidine-1-carboxylic acid 1-tert-butyl ester.

A solution of 2-(S)-carbamoyl-4-oxo-piperidine-1-carboxylic acid 1-tert butyl ester (3.51 g, 14.48 mmol), tert-butyl cyanoacetate (2.04 g, 14.48 mmol), sulphur (0.464 g, 14.48 mmol) and diisopropyl ethylamine (2.5 mL,

14.48 mmol) in methanol (20 mL) was heated 16 hours at 40°C under N<sub>2</sub>. The volatiles were removed in vacuo and the residue was purified using column chromatography (SiO<sub>2</sub>, Flash 40, petroleum ether/ethyl acetate 3:1) to give 1.33 g (23%) of a mixture 2-amino-5-(*S*)-carbamoyl-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,6-di-carboxylic acid di-*tert*-butyl ester and 2-amino-7-(*R*)-carbamoyl-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,6-di-carboxylic acid di-*tert*-butyl ester isomers.

0.5 g (1.25 mmol) of the above mixture was dissolved dichloromethane
(10 mL) and imidazole-1-yl-oxo-acetic acid *tert*-butyl ester (0.74 g, 3.77 mmol) and triethylamine (0.525 mL, 3.77 mmol) was added. The reaction mixture was stirred for 16 hours at room temperature before the volatiles were removed in vacuo. The residue was purified by column chromatography (SiO<sub>2</sub>, Flash 40, petroleum ether/ethyl acetate (4:1)) too
give 75 mg (11%) of 2-(*tert*-butoxyoxalyl-amino)-7-(*R*)-carbamoyl-4,7-dithydro-5*H*-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester. This was dissolved in a mixture of trifluoacetic acid/dichloromethane (1:1) (10 mL) and stirred for 16 hours at room temperature before the solvent was removed in vacuo. The residue was recrystallized from methanol to give 24 mg (39%) of the title compound.

LC-MS;  $R_t = 1.56$  min, m/z:  $314 [M+H]^{+}$ Calculated for  $C_{11}H_{11}N_3O_6S$ ,  $0.25xC_2HF_3O_2$ ,  $0.75xH_2O$ C, 38.88 %; H, 3.62 %; N, 11.83 %; Found: C, 38.92 %; H, 3.92 %; N, 11.81 %.

#### **EXAMPLE 113**

30 2-(Oxalyl-amino)-5-(S)-(2-oxo-tetrahydro-thiophen-3-ylcarbamoyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

A solution of 2-amino-4,7-dihydro-5*H*-thieno[2,3-c]pyridine-3,5-(*S*),6-tricarboxylic acid 3,6-di-*tert*-butyl ester (0.30 g, 0.75 mmol) and triethylamine (0.21 mL, 1.51 mmol) in tetrahydrofuran (10 mL) was cooled to -20°C before isobutyl chloroformate (0.103 mL, 0.75 mmol) was added. The reaction mixture was stirred 15 min at -20°C before homocystein hydrochloride (116 mg, 0.75 mmol) was added. The cooling bath was removed and the reaction mixture was left for 16 hours at room temperature. The solvent was removed in vacuo and the residue was subjected to column chromatography (SiO<sub>2</sub>, Flash 40, heptane/ethyl acetate 2:1) to give 212 mg (56%) of 2-amino-5-(*S*)-(2-oxo-tetrahydro-thiophen-3-ylcarbamoyl)-4,7-dihydro-5*H*-thieno[2,3-*c*]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester

A solution of 2-amino-5-(*S*)-(2-oxo-tetrahydro-thiophen-3-ylcarbamoyl)4,7-dihydro-5*H*-thieno[2,3-*c*]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester (200 mg, 0.40 mmol), imidazole-1-yl-oxo-acetic acid *tert*-butyl ester (235 mg, 1.20 mmol) and triethylamine (168 μL, 1.20 mmol) in dichloromethane (10 mL) was stirred for 16 hours at room temperature before the solvent was removed <u>in vacuo</u>. The residue was purified by column chromatography (SiO<sub>2</sub>, Flash 40, heptane/ethyl acetate 2:1) to give 250 mg (100%) of 2-(*tert*-butoxyoxalyl-amino)-5-(*S*)-(2-oxotetrahydro-thiophen-3-ylcarbamoyl)-4,7-dihydro-5*H*-thieno[2,3-*c*]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester.

This was dissolved in a mixture of trifluoroacetic acid/dichloromethane
(1:1) (3 mL) and stirred for 16 hours at room temperature before diethyl
ether (6 mL) was added. The precipitate was filtered off and washed with
diethyl ether to give 172 mg (81%) of the title compound as a solid
trifluoroacetate.

LC-MS;  $R_t = 0.41 \text{ min, m/z: } 414 \text{ [M+H]}^{+}$ 

30 Calculated for C<sub>15</sub>H<sub>15</sub>N<sub>3</sub>O<sub>7</sub>S<sub>2</sub>, 1.5xC<sub>2</sub>HF<sub>3</sub>O<sub>2</sub>, H<sub>2</sub>O; C, 35.88 %; H, 3.10 %; N, 6.97 %; Found: C, 35.91 %; H, 3.54 %; N, 6.97 %.

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#### **EXAMPLE 114**

2-(Oxalyl-amino)-5-(S)-phenylcarbamoyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

A solution of 2-amino-4,5,6,7-tetrahydro-thieno[2,3-c]-pyridine-3,5-(S),6tricarboxylic acid 3,5-di-tert-butyl ester (300 mg, 0.75 mmol) and triethylamine (210 µL, 1.51 mmol) in tetrahydrofuran (10 mL) was cooled to -20°C before isobutylchloroformate (103 mg, 0.75 mmol) was introduced. The reaction mixture was stirred for 20 min before aniline (70 mg, 0.75 mmol) was added. The cooling bath was removed and the reaction was left for 16 hours at room temperature before the solvent was removed in vacuo. The residue was partitioned between ethyl acetate (50 mL) and water (50 mL). The organic phase was dried (MgSO<sub>4</sub>) and the solvent was removed in vacuo. The residue was dissolved in dichloromethane (10 mL) and imidazole-1-yl-oxo-acetic acid tert-butyl ester (443 mg, 2.26 mmol) and triethylamine (315 µL, 2.26 mmol) was added. The reaction mixture was stirred 16 hours at room temperature before the solvent was removed in vacuo. The residue was purified by column chromatography (SiO2, Flash 40, heptane/ethyl acetate (3:1) to give 250 mg 2-(tert-butoxyoxalyl-amino)-5-(S)-phenylcarbamoyl-4,7dihydro-5*H*-thieno[2,3-*c*]pyridine-3,6-dicarboxylic acid di-*tert*-butyl ester.

2-(tert-Butoxyoxalyl-amino)-5-(S)-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester was dissolved in a mixture of trifluoroacetic acid/dichloromethane (1:1) (3 mL) and stirred for 16 hours at room temperature before diethyl ether (6 mL) was added. The precipitate was filtered off and washed with diethyl ether to give 155
 mg (41%) of the title compound as a solid trifluoroacetate. LC-MS; R<sub>t</sub> = 0.86 min, m/z: 390 [M+H]<sup>+</sup>

Calculated for  $C_{17}H_{15}N_3O_6S$ ,  $1.5xC_2HF_3O_2$ ,  $H_2O$ ; C, 41.53 %; H, 3.22 %; N, 7.26 %; Found: C, 41.77 %; H, 3.29 %; N, 7.28 %.

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#### **EXAMPLE 115**

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2-(Oxalyl-amino)-7-(R)-phenylcarbamoyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

To a solution of 2-(*S*)-4-oxo-piperidine-1,2-dicarboxylic acid 1-*tert* butyl ester (2.06 g, 8.47 mmol) and triethylamine (1.42 mL, 10.16 mmol) in tetrahydrofuran (20 mL) cooled to -20°C was added isobutylchloroformate (1.39 g, 10.16 mmol) and the mixture was stirred for 10 min at -20°C before aniline (946 mg, 10.16 mmol) was added. The cooling bath was removed and the reaction mixture was stirred for 16 hours at room temperature before the solvent was removed <u>in vacuo</u>. The residue was divided between water (50 mL) and ethyl acetate (50 mL). The organic phase was washed with saturated sodium chloride (25 mL) and dried (MgSO<sub>4</sub>). After filtration and concentration <u>in vacuo</u> the residue was purified using column chromatography (SiO<sub>2</sub>, Flash 40, petroleum ether/ethyl acetate 5:1) to give 1.3 g (48%) of 4-oxo-2-(*S*)-phenyl-carbamoyl-piperidine-1-carboxylic acid *tert*-butyl ester.

A solution of 4-oxo-2-(S)-phenylcarbamoyl-piperidine-1-carboxylic acid *tert*-butyl ester (1.3 g, 4.08 mmol), *tert*-butylcyanoacetate (0.58 g, 4.08 mmol), sulphur (0.133 g, 4.08 mmol) and diisopropyl ethylamine (0.7 mL,

4.08 mmol) in methanol (10 mL) was heated under nitrogen to 40 °C for 16 hours before the solvent was removed in vacuo. The residue was subjected to column chromatography (SiO<sub>2</sub>, Flash 40, petroleum ether/ethyl acetate 6:1) to give 0.70 g (36%) of a mixture of 2-amino-5-(S)phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-di-carboxylic 5 acid di-tert-butyl ester and 2-amino-7-(R)-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-di-carboxylic acid di-tert-butyl ester isomers. The above mixture was dissolved in dichloromethane (20 mL) and imidazole-1-yl-oxo-acetic acid tert-butyl ester (872 mg, 4.44 mmol) and triethylamine (618 µL, 4.44 mmol) was added. The reaction mixture was 10 stirred 16 hours before the solvent was removed in vacuo and the residue was subjected to column chromatography (SiO<sub>2</sub>, Flash 40, petroleum ether/ethyl acetate 5:1) to give 0.50 g (56%) as a mixture of 2-(tertbutoxyoxalyl-amino)-5-(S)-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3c]pyridine-3,6-di-carboxylic acid di-tert-butyl ester and 2-(tert-butoxyoxalyl-15 amino)-7-(R)-phenylcarbamoyl-4,7-dihydro-5H-thieno[2,3-c]pyridine-3,6-dicarboxylic acid di-tert-butyl ester

300 mg of the mixture was dissolved in a mixture of trifluoacetic
20 acid/dichloro-methane (1:1) (6.0 mL) and the solution was stirred for 16 hours at room temperature before the solvent was removed in vacuo. The residue was purified on preparative HPLC to give 70 mg (34%) of the title compound as a solid trifluoroacetate.

25 LC-MS;  $R_t = 0.95$  min, m/z: 390 [M+H]<sup>+</sup> Calculated for  $C_{17}H_{15}N_3O_6S$ ,  $C_2HF_3O_2$ ,  $H_2O$ ; C, 43.77 %; H, 3.48 %; N, 8.06 %; Found: C, 43.92 %; H, 3.44 %; N, 7.97 %.

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EXAMPLE 116



5-(R),7-(R)-Bis-benzyloxymethyl-2-(oxalyl-amino)-4,5,6,7-tetrahydrothieno[2,3-c]pyridine-3-carboxylic acid

Benzyloxyacetaldehyde (0.90 g; 6.0 mmol) and dimethyl (2oxomethyl)phosphonate (1.0 g; 6.0 mmol) were dissolved in a mixture of tetrahydrofuran (25 ml) and water (20 ml). 1N Aqueous potassium hydroxide (6 ml) was added and the mixture was stirred for 30 min. Dichloromethane (50 ml) was added and the organic phase was

separated, dried (MgSO<sub>4</sub>) and evaporated in vacuo leaving 5-10 benzyloxypent-3-en-2-one.

<sup>1</sup>H-NMR: 2.25 (s, 3H); 4.19 (dd, 2H); 4.55 (s, 2H); 6.34 (dt; 1H); 6.70 (dt, 1H); 7.26 (m, 5H).

5-benzyloxypent-3-en-2-one was dissolved in methanol (5 ml) and ammonium acetate (13 mmol, 1.03 g) was mixted together with 15 benzyloxyacetaldehyde (1.8 g; 12 mmol) and acetic acid (0.69 ml) and the mixture was stirred for 2 days. The solvent was removed in vacuo and the residue was chromatographed on silica using gradient elution from 100 % dichloromethane to 100 % ethyl acetate. A fraction (411 mg) contained 20 (according to LC-MS; m/z 340.4) 2,5-di(benzyloxymethyl)-4-piperidone in

an impure state was isolated. The crude mixture was dissolved in ethanol (3 ml) and tert-butylcyanoacetate (400 mg), sulfur (100 mg) and triethylamine was added and the mixture was stirred at room temperature overnight. The mixture was filtered and the solvent removed in vacuo. The

residue was chromatographed on silica in a mixture of dichloromethane/(7% of 25% aqueous ammonia in ethanol) (40:1), which afforded 0.14 g of 2-amino-5-(R),7-(R)-bis-benzyloxymethyl-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester as an oil.

LC-MS: R<sub>t</sub>: 6.03 min; m/z: 495.2 [M+H]<sup>+</sup>

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2-amino-5-(*R*),7-(*R*)-Bis-benzyloxymethyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (0.14 g; 0.28 mmol) was dissolved in dichloromethane (5 ml) and treated with imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.1 g; 0.5 mmol) and triethylamine (70 μl; 0.5 mmol), and stirred overnight, washed with water, dried (MgSO<sub>4</sub>) and the solvent removed <u>in vacuo</u>. The residue was chromatographed on silica using ethyl acetate/dichloromethane (1:3) as eluent. The residue was treated with trifluoroacetic acid (0.5 ml) in dichloromethane (0.5 ml) and stirred for 4 hours. Evaporation of the solvent <u>in vacuo</u> afforded 37 mg of the <u>title compound</u>.

LC-MS: R<sub>t</sub>: 4.74 min; m/z: 511.4 [M+H]<sup>+</sup>.

#### **EXAMPLE 117**

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6-Benzyl-2-(oxalyl-amino)-5-(1,1,3-trioxo-1,3-dihydro-1,6-benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid

1-Benzyl-4-oxo-piperidine-2-carboxylic acid ethyl ester (2.9 g; 11.1 mmol) (prepared in a similar way as described in "GENERAL CHIRAL SYNTHESIS" for 4-oxo-1-((S)-1-phenyl-ethyl)-piperidine-(R)-2-carboxylic acid ethyl ester using benzylamine instead of 1-(S)-phenethylamine) was dissolved in abs. ethanol (50 ml) and sulfur (0.35 g, 11.1 mmol), triethylamine (1.6 ml, 11.1 mmol), and *tert*-butylcyanoacetate (1.7 g, 11.1 mmol) were added and the mixture was stirred 2 days at room temperature. The solvent was removed <u>in vacuo</u> and the residue was chromatographed on silica using a mixture of ethyl actetate/heptane (1.4) as eleuent leaving a mixture (700 mg; 1:1 based on NMR) of 2-amino-6-benzyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3,7-dicarboxylic acid 3-*tert*-butyl ester-7-ethyl ester and 2-amino-6-benzyl-4,5,6,7-tetrahydro-

thieno[2,3-c]pyridine-3,7-dicarboxylic acid 3-tert-butyl ester 5-ethyl ester which was used in the next step without separation. To this mixture was added tetrahydrofuran (5 ml) and lithium borohydride (1.1 ml of a 2M solution in tetrahydrofuran) and the mixture was stirred 18 hours. More lithium borohydride (5.0 ml of a 2M solution in tetrahydrofuran) was added and the mixture stirred for an additiona 4 days. Ethyl acetate (10 ml) was added dropwise and after 1 hour the mixture was poured onto water (100 ml) and extracted with dichloromethane (2 x 100 ml) and chromatographed on silica (using ethylacetate/heptane 1:1 as eluent), which afforded a mixture of 2-amino-6-benzyl-7-hydroxymethyl-4,5,6,7tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid tert-butyl ester and 2amino-6-benzyl-5-hydroxymethyl-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (in total 187 mg). To this mixture was added dry tetrahydrofuran (10 ml), 2,3-dihydro-1,2-benzisothiazol-3-one-15 1,1-dioxide (100 mg; 0.55 mmol), triphenylphosphine (144 mg 0.55 mmol) and the mixture was cooled with ice. Diethyl azodicarboxylate (86 µl) was added and the mixture was stirred overnight at room temperature. The solvent was evaporated in vacuo and the residue was chromatographed on silica using a mixture of ethyl acetate/heptane (1:1) as eluent leaving 94 mg of 2-amino-6-benzyl-5-(1,1,3-trioxo-1,3-dihydro-1,6-20 benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3carboxylic acid tert-butyl ester. <sup>1</sup>H-NMR: (CDCl3): 1.52 (s, 9H); 2.75 (dd, 1H); 2.90 (dd, 1H); 3.55 (d, 1H); 3.72 (m, 4H); 3.94 (d, 1H); 4.12 (d, 1H); 5.97 (s, 2H); 7.14-7.37 (m, 5H); 25 7.80-8.03 (m, 4H).

2-Amino-6-benzyl-5-(1,1,3-trioxo-1,3-dihydro-1,6-benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (94 mg; 0.17 mmol) was dissolved in dichloromethane (5 ml) and treated with imidazol-1-yl-oxo-acetic acid *tert*-butyl ester (0.07 g; 0.3 mmol) and triethylamine (49 μl; 0.3 mmol), and stirred overnight,-washed-with water, 1N aqueous citric acid, dried (MgSO<sub>4</sub>) and the solvent

LC-MS: Rt 5.47 min, m/z: 540.4 [M+H]

removed <u>in vacuo</u> leaving 104 mg of 2-(*tert*-butoxyoxalyl-amino)-6-benzyl-5-(1,1,3-trioxo-1,3-dihydro-1,6-benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester as an oil. LC-MS:  $R_t$ : 5.50 min, m/z: 668.6 [M+H]<sup>+</sup>

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2-(*tert*-Butoxyoxalyl-amino)-6-benzyl-5-(1,1,3-trioxo-1,3-dihydro-1,6-benzo[d]isothiazol-2-ylmethyl)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid *tert*-butyl ester (100 mg; 0.15 mmol) was treated with trifluoroacetic acid (1 m) in dichloromethane (4 ml) and stirred for 2 days.

Evaporation of the solvent in vacuo afforded 90 mg of the <u>title compound</u> as a solid trifluoroacetate.

Calc. for  $C_{25}H_{21}N_3O_8S_2$ ,  $1.5xC_2HF_3O_2$ ,  $0.5xH_2O$  C, 45.72%; H, 3.22%; N, 5.71%. Found:

C, 45.48%; H, 3.46%; N, 5.72%

15 LC-MS: R<sub>t</sub>: 4.16 min; m/z: 556.2 [M+H]<sup>+</sup>

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## **EXAMPLE 118**

# Crystallisation of protein and protein-inhibitor complexes

Co-crystallization of PTP1B with inhibitors;

A 6-10 mg/ml preparation of PTP1B in 10 mM Tris pH 7.5, 25 mM NaCl, 0.2 mM EDTA and 3 mM DTT, was used for crystallization. Crystals were grown by the sitting as well as the hanging drop vapor diffusion methods. A 1:10 (PTP1B:inhibitor) molar ratio mixture was prepared at least one hour prior to crystallization. Two μl of PTP1B-inhibitor solution was mixed with 2 μl reservoir solution consisting of: 0.1 M Hepes buffer pH 7.5, 0.3-0.4 M Na-acetate or Mg-acetate, 12-16% Peg 8000 and/or 4% glycerol. The reservoir volume was 1 ml. Crystals grew to the size of 0.3-0.6X0.1-0.3X0.1-0.3 mm over 2-3 days.

Data collection.

All crystal data collections were performed at 100 K. The following cryo conditions were used: to the hanging or sitting drop 3  $\mu$ l of 50% glycerol (containing 0.5 mmol inhibitor) were added. The crystal was removed from the drop after 5-30 min. and transferred to 50% glycerol (containing 0.5 mmol inhibitor) and rapidly flash frozen.

Data were collected using a mar345 image plate either at the MAX-lab synchrotron facilities in Lund (Sweden) or in-house equipped with a rotating anode (RU300) and Osmic multilayer mirror system. Typically a 1° oscillation was used for 60 images data sets in the resolution range 2.7-1.8 Å were obtained. The space group was determined to be P3121 for all crystals used.

#### Refinements.

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As P3121 contains a polar axis and, thus, possesses more than one indexing possibility, a molecular replacement solution using Amore [ref] solution was found prior to the refinements. A high resolution PTP1B structure was used as a starting model, with ligand and water molecules omitted from the structure. All refinements were performed with 'Xplor. v. 3.851 [MSI]. Interchanging cycles of model building using X-build [MSI] and refinement were performed. The 2Fo-Fc maps were inspected by the use of X-ligand [MSI] at a 1.3 sigma level for densities that could correspond to the structures of the inhibitors. In all cases a well-suited inhibitor electron density was identified in the active site pocket, see figures 1-4. No other densities were identified to fit the inhibitors. Water molecules were inserted using the X-solvate program [MSI].



Coupling of 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid to Epoxy-activated Sepharose 6B.

This example describes the preparation of an immobilized compound suited for affinity chromatographic purification of PTPases (eg PTP1B or T-cell PTP).

3.5 g Epoxy-activated Sepharose 6B (Pharmacia Biotech) was prepared for coupling according to the manufacturers directions, and divided into 3 portions (3 x 8 ml gel-suspension, corresponding to 4 ml drained gel each).

8 ml portions of 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid dissolved to 10, 1 and 0.1 mM in a 0.2 M sodium carbonate coupling buffer pH 9 were mixed with the gel suspensions and agitated gently overnight at room temperature.

Exces ligand was washed away, the remaining active groups were blocked and the product was washed extensively at alternating pH, all according to the the manufacturers directions.

The products were stored refrigerated in 0.1 M acetate pH 4.0 containing 0.5 M sodium chloride.

Significant inhibition of PTP1B was demonstrated in the 20 µmole ligand/ml gel preparation, when diluted to 1 µl drained gel/ml.

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#### **EXAMPLE 120**

Affinity purification of PTP1B using the compound 2-(oxalyl-amino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid coupled to Epoxy-activated Sepharose 6B.

This example describes the affinity chromatographic purification of a PTPase.

2 ml of the product with 20 µmole ligand/ml described in example 55 was loaded into a 1.6 cm diameter column and equilibrated with a buffer (buffer A) containing

20 mM L-histidine

5 1 mM EDTA

7 mM Mercaptoethanol

100 mM Sodium chloride

and adjusted to pH 6.2 with 1 M HCI.

1.5 mg conventionally purified PTP1B in 5 ml buffer A, was applied to the column at 0.5 ml/min followed by a wash with 10 ml buffer A.

UV absorbing material without PTPase activity, corresponding to approx.

10 % of the totally applied material, passed through the column.

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The flow direction was reversed, the flow increased to 2 ml/min and linear gradient elution started with a combined salt and pH gradient for 20 minutes using buffer B containing

20 mM L-histidine

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1 mM EDTA

7 mM Mercaptoethanol

1 M Sodium chloride

and adjusted to pH 9.0 with 1 M NaOH.

25 Maximum elution took place at approx. 32 % buffer B (0.39 M NaCl and pH 6.8) in a broad peak.

The total activity yield in the elution peak was 70 %, and the specific activity of the enzyme was improved by a factor 1.4.

## **EXAMPLE 121**

Use of compounds of the invention to identify substrates that are specifically dephosphorylated by PTPases that are inhibited by the compounds of the invention or by other PTPases.

The compounds of the invention are unique tools for identification of cellular substrates of the PTPases that are inhibited by the compounds of the invention. Substrates are herein defined as cellular proteins that (i) are phosphorylated on tyrosine residues, (ii) are dephosphorylated by PTPases that are inhibited by compounds of the invention or by other PTPases. If said substrates are dephosphorylated by PTPases that are inhibited by compounds of the invention, administration of the compounds of the invention will result in partial or total prevention of dephosphorylation of said substrates. As a result, a concomitant prolonged or increased activation may be observed of the signal transduction pathway (for definition, *vide infra*) in which said substrate is involved. Non-limiting examples of substrates are: the insulin receptor  $\beta$  subunit, IRS-1, IRS-2, IRS-3, IRS-4, JAK1, JAK2, shc-2, grb-2 (Hunter, *Cell 100*: 113-127 (2000)).

Importantly, the compounds of the invention can also be used to identify novel substrates. When the compounds of the invention have been used to identify the substrates of the PTPases that are inhibited by the compounds of the invention, a person skilled in the arts will be able to use this knowledge to establish animal models that will reflect a human condition or disease in which a compound of the invention will be indicated. Non-limiting example of the usefulness of said compounds of the invention will be in the following disease areas: diabetes, obesity cancer and conditions with unwarranted platelet aggregation.

To identify the substrates of the PTPases that are inhibited by the substrates of the invention the following methods may be employed.

Whole animals and/or primary cells and/or cell lines that represent the

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target organ or tissue may be used for these experiments. Non-limiting examples of animals are: ob/ob mice (worldwide web @ jax.org); db/db mice: Zucker obese rats. Non-limiting examples of target tissues or organs are: skeletal muscle, liver, adipose tissue, pancreas, the spleen, the bone marrow. Non-limiting examples of cell lines are: Chinese hamster ovary (CHO) cells (CHO-K1 – American Type Culture Collection (ATCC) Number CCL-61), Baby Hamster Kidney (BHK) cells (ATCC Number CRL-1632), HepG2 cells (ATCC Number HB-8065), C2C12 cells (ATCC Number CRL-1772), L6 cells (ATCC Number CRL-1458), RD cells (ATCC Number CCL-136). Said cells can either be unmanipulated or transfected transiently or permanently with plasmid vectors that encode proteins or substrates. Non-limiting example of a plasmid that allows expression in mammalian cells are: pcDNA1 and pcDNA3 (worldwode web @ invitrogen.com). Non-limiting examples of proteins or substrates that are transfected into said cell lines are: the insulin receptor, the IGF-I receptor, the EGF-R receptor, the PDGF receptor, IRS-1, IRS-2, IRS-3, IRS-4, p56Lck; Jak1, Jak2 (Hunter, supra).

The analysis consists of the following steps:

20 (A) stimulation of signal transduction pathways with and without the presence of the compounds of the invention. Signal transduction pathways are herein defined as a series of cellular processes that are initiated by a triggering event (such as stimulation of a tissue or cell by a hormone and/or a cytokine and/or cell-cell interaction and/or cell-cell substratum interaction) leading to various cellular effects including 25 metabolic effects, cell differentiation and cell proliferation (Hunter, supra). Non-limiting examples of signal transduction pathways include: the insulin signaling pathway; the leptin signalling pathway; thrombin signalling pathway; the erythropoietin signaling pathway; the epidermal growth factor signaling pathway. Non-limiting examples of the effects of stimulating 30 signal transduction pathways: glucose uptake; glycogen synthesis; cell proliferation; cell differentiation; platelet aggregation-

(B) Analysis and identification of substrates that show increased (or decreased) phosphorylation on tyrosine residues after administration of the compound of the invention in comparison with controls that did not receive the compound.

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## Step A. Stimulation of signal transduction pathways.

As a non-limiting example, insulin (concentration range: 0.1 to 100 nM. final concentration) is administered to primary hepatocytes in tissue culture plates. The compounds of the invention (concentration range: 10 nM to 100 μM) are administered to half of the plates, with the other plates acting as controls. The plates are incubated at 37 °C for various time periods: Typically for 0, 1, 2, 5, 15, 30 and 60 mins. Following this stimulation, the plates are treated as follows: The medium is rapidly aspirated and the cells washed twice with ice-cold PBS. Two milliliters of ice-cold lysis buffer (see below) is added and the plates are placed on ice for 2 minutes after which the cells are scraped off using a cell scraper ('rubber policeman'). The lysates are placed at 4 °C at a rotary shakerr. Dithiotreitol is added to a final concentration of 10 mM, and the lysates are centrifuged at 20,000 r.p.m.. Aliquots of the supernatants, i.e. lysates, are stored at -80 °C until further use.

Lysis buffer – for a total of 20 ml add the following

0.8 ml of 500 mM Tris-Cl, pH 7.4

0.2 ml of 100 mM EDTA

2.0 ml of 1 M NaCl

2.0 ml of 10 % (vol/vol) Triton X-100 25  $80 \mu l$  of 250 mM PMSF

2 µl of 10 mg/ml aprotinin

20 µl of 1 mg/ml leupeptin

5 mM 100 mM iodoacetate

11.88 ml demineralized water 30

> Step B. Analysis and identification of substrates that show increased (or decreased) phosphorylation on tyrosine residues after administration of the compound of the invention in comparison with controls that did not receive the compound.

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skilled in the art.



As a non-limiting example, said lysates are subjected to twodimensional polyacrylamide gel electrophoresis (2-D PAGE) followed by detection of proteins that are phosphorylated on tyrosine residues (pTyr) by western blotting, techniques well-known to those skilled in the art (Marcus et al. Electrophoresis 21: 2622-2636 (2000)). Proteins that show increased (or decreased) pTyr are identified by comparing the western blots made from said lysates derived from said hepatocytes treated with both insulin and the compounds of the invention with said control lysates derived from said hepatocytes that were treated with insulin only. Increased pTyr of a protein shows that the said protein is regulated by the PTPase or PTPases that are inhibited by the compounds of the invention. Said protein may either be a direct substrate of the PTPase or PTPases that are inhibited by the compounds of the invention or the substrate of other PTPase(s) which activity is regulated by the PTPase or PTPases that are inhibited by the compounds of the invention. Decreased pTyr of a protein shows that said protein is the substrate of other PTPase(s) that is/are activated, directly or indirectly, by the PTPase or PTPases that are inhibited by the compounds of the invention. Having identified and visualized proteins, i.e. substrates, that show changed pTyr levels, the spots are cut out, digested with trypsin and analyzed by matrix assisted laser desorption/ionization-time of flight-mass spectrometry (MALDI-TOF-MS) (Marcus et al., supra). To identify the nature of said substrate with changed pTyr levels the obtained mass fingerprints are analyzed as described by Marcus et al. (supra) or other methods well-known to those

Said substrate can either be an already described protein or a novel protein. In both cases, the identification may be followed by cDNA cloning procedures with the aim of obtaining a full-length clone corresponding to said substrate using standard techniques well-known to those skilled in the art (Ausubel, F. M., *et al.* (ED.). Short Protocols in Molecular Biology, 2<sup>nd</sup> ed, John Wiley and Sons, inc., New York, ISBN 0-471-57735-9-(1992)). Said-full-length clone may be expressed as recombinant proteins in prokaryotic or eukaryotic expression systems well-known to those skilled in the art (worldwide web @ invitrogen.com;

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worldwide web @ stratagene.com, worldwide web @ promega.com), and the function of said substrate may in turn be studied both at the biochemical and cellular levels. Further, said recombinant proteins may further be used as an antigen to produce either polyclonal or monoclonal antibodies using techniques well-known to those skilled in the art. As a non-limiting example, with said full-length clone, said antibodies, and the compounds of the invention at hand, those skilled in the art will be able to study the tissue distribution and expression levels of said substrates in normal animals and animal models of diseases, such as diabetes, obesity, cancer and disturbances of platelet aggregation. A person skilled in the art will be able to use this knowledge to establish animal models or use already established animal models that will reflect a human condition or disease in which a compound of the invention will be indicated. Non-limiting example of the usefulness of said compounds of the invention will be in the following disease areas: diabetes, obesity, cancer and conditions

### **EXAMPLE 122**

with unwarranted platelet aggregation.

Identification of substrates that are dephosphorylated by PTPases that are inhibited by the compounds of the invention

The analysis consists of the following steps: (A) preparation of hyperphosphorylated substrates; (B) identification of said substrates that are dephosphorylated by PTPases that are dephosphorylated by compounds of the invention.

To identify the substrates of the PTPases that are inhibited by the compounds of the invention the following method may be employed. Primary cells and/or cell lines that represent the target organ or tissue may be used for these experiments. Non-limiting examples of target tissues or organs are: skeletal muscle, liver, adipose tissue, pancreas, the spleen, the bone marrow. Non-limiting examples of cell lines are: Chinese hamster ovary (CHO) cells (CHO-K1 – American Type Culture Collection (ATCC) Number CCL-61), Baby Hamster Kidney (BHK) cells (ATCC Number CRL-1632), HepG2 cells (ATCC Number HB-8065), C2C12 cells (ATCC

Number CRL-1772), L6 cells (ATCC Number CRL-1458), RD cells (ATCC Number CCL-136). Said cells can either be unmanipulated or transfected transiently or permanently with plasmid vectors that encode proteins or substrates. Non-limiting example of a plasmids that allow expression in mammalian cells are: pcDNA1 and pcDNA3 (worldwide web @ invitrogen.com). Non-limiting examples of proteins or substrates that are transfected into said cell lines are: the insulin receptor, the IGF-I receptor, the EGF-R receptor, the PDGF receptor, IRS-1, IRS-2, IRS-3, IRS-4, p56Lck; Jak1, Jak2 (Hunter, *supra*).

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### Step A

Said primary cells, tissues or cell lines are exposed to a general inhibitor of PTPases. This treatment results in induction of hyperphosphorylation of a multitude of cellular substrates. A non-limiting example of a general PTPase inhibitor is bisperoxovanadium 1,10 phenanthroline (bpV(phen)) (Posner *et al. J. Biol. Chem.* 269: 4596-4604 (1994)).

A non-limiting example of a hyperphosphorylation protocol: CHO cells that stably overexpress the insulin receptor are grown in 15 cm Petri dishes to 80-90 percent confluence (using F-12 medium with 10 percent fetal calf serum). The culture medium is replaced with medium that does not contain calf serum and are grown for additional 2 hrs at 37 °C. The plates are washed twice with phosphate buffered saline (PBS) and incubated for further 2 hours with 100 µM bpV(phen) and 100 nM insulin (Novo Nordisk) (final assay concentrations). Following this stimulation the plates are treated as follows: The medium is rapidly aspirated and the cells washed twice with ice-cold PBS. Two milliliters of ice-cold lysis buffer (see below) is added and the plates are placed on ice for 2 minutes after . which the cells are scraped off using a cell scraper ('rubber policeman'). The lysates are placed at 4 °C at a rotary shaker for 1 hour. Dithiotreitol is added to a final concentration of 10 mM, and the lysates are centrifuged for 10 minutes at 20,000-r.p.m.- Aliquots of the supernatants, i.e. lysates, are stored at -80 °C until further use.

Lysis buffer – for a total of 20 ml add the following:

0.8 ml of 500 mM Tris-Cl, pH 7.4

0.2 ml of 100 mM EDTA

2.0 ml of 1 M NaCl

5 2.0 ml of 10 % (vol/vol) Triton X-100

 $80 \mu l$  of 250 mM PMSF

2 μl of 10 mg/ml aprotinin

20 μl of 1 mg/ml leupeptin

5 mM 100 mM iodoacetate

10 11.88 ml demineralized water

### Step B

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For these studies both novel and known PTPases may be used. The PTPases may be either isolated using the compounds of the invention as described in Example 120 or recombinant proteins. Non-limiting examples of known PTPases that are inhibited by compounds of the invention are PTP1B and TC-PTP. The cDNA for these PTPases are inserted in prokaryotic expression vectors and are expressed in *E. coli*. An overnight culture is diluted 1:25 into a total volume of 2 liters of SOB medium and grown at 37 °C for 3 hours. Isopropyl  $\beta$ -D-thiogalactoside (IPTG) is added to a final concentration of 0.1 mM, and the incubation is continued at room temperature for 3 hrs. The fusion proteins are purified according to the manufacturer's instructions (Amersham Pharmacia Biotech).

25 Aliquots of said lysates (60 μl) are mixed with said PTPase that is inhibited by said compound of the invention and incubated on ice for 1, 10, and 30 minutes. At each time point, 20 μl aliquots are removed and mixed with SDS loading buffer (20% (v/v) glycerol, 3% (w/v) SDS, 3% (v/v) 2-mercaptoethanol, 10 mM EDTA, 0.05% (w/v) bromphenol blue), heated at 100 °C for 2 minutes and stored at – 20 °C until use. Control lysates without addition of PTPase are treated identically.

As a non-limiting example, said lysates are subjected to two-dimensional polyacrylamide gel electrophoresis (2-D PAGE) followed by detection of proteins that are phosphorylated on tyrosine residues (pTyr) by western blotting, techniques well-known to those skilled in the art (Marcus *et al.* 

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Electrophoresis 21: 2622-2636 (2000)). Proteins that show decreased pTyr are identified by comparing the western blots made from said lysates treated with said PTPase with said control lysates. Decreased pTyr of a protein shows that the said protein is a substrate of the PTPase or PTPases that are inhibited by the compounds of the invention. Having identified and visualized proteins, i.e. substrates, that show decreased pTyr levels, the spots are cut out, digested with trypsin and analyzed by matrix assisted laser desorption/ionization-time of flight-mass spectrometry (MALDI-TOF-MS) (Marcus et al., supra). To identify the nature of said substrate with decreased pTyr levels the obtained mass fingerprints are analyzed as described by Marcus et al. (supra) or other methods well-known to those skilled in the art.

Said substrate can either be an already described protein or a novel protein. In both cases, the identification may be followed by cDNA cloning procedures with the aim of obtaining a full-length clone corresponding to said substrate using standard techniques well-know to those skilled in the art (Ausubel, F. M., *et al.* (ED.). Short protocols in molecular biology, 2<sup>nd</sup> ed, John Wiley and sons, inc., New York, ISBN 0-471-57735-9 (1992)). Further use of the knowledge include analysis in animal models as described in Example 59

#### **EXAMPLE 123**

## Analysis for blood glucose lowering effects

The compounds of the invention are tested for blood glucose lowering effects in diabetic, obese female *ob/ob* mice. The mice are of similar age and body weights and they are randomized into groups of ten mice. They have free access to food and water during the experiment. The compounds are administered by either by gavage, subcutaneous, intravenous or intraperitoneal injections. The control group receives the same volume of vehicle as the mice that receive the compounds. Non-limiting examples of dose-range: 0.1, 0.3, 1.0, 3.0,10, 30, 100 mg per kg body weight. The blood-glucose-levels are measured two times before administration of the compounds of the invention and vehicle (to the control group). After administration of the compound, the blood glucose

levels are measured at the following time points: 1, 2, 4, 6, and 8 hours. A positive response is defined either as (i) a more than 25 percent reduction in blood glucose levels in the group receiving the compound of the invention compared to the group receiving the vehicle at any time point or (ii) statistically significant (i.e. p<0.05) reduction in the area under the blood glucose curve during the whole period (i.e. 8 hrs) in the group treated with the compounds of the invention compared to the group receiving the vehicle.

All documents cited herein are incorporated by reference in their entirety.

In case of conflict in definitions, the present definitions control.



# TABLE A

Table of the orthogonal three dimensional coordinates in Ångstroms and B factors (Ų) for Protein Tyrosine Phosphatase 1B complexed with 2-(oxalylamino)-4,5,6,7-tetrahydro-thieno[2,3-c]pyridine-3-carboxylic acid.

No	Amin	o acid	X	Υ	Z	В			
1	GLU	N	69.819		14.736		20.949		51.88
2	GLU	CA	69.381		16.070		20.592		50.12
3	GLU	С	68.816		16.123		19.177		50.96
4	GLU	O	69.477		15.855		18.147		47.57
5	GLU	СВ	70.340		17.247		20.871		48.87
6	GLU	CG	69.694		18.607		20.512		43.78
7	GLU	CD	68.658		19.051		21.547		100.00
8	GLU	OE1	68.838		19.978		22.327		100.00
9	GLU	OE2	67.553		18.331		21.549	•	100.00
10	MET	N	67.550°		16.476		19.190		38.96
11	MET	CA	66.810	6 a	16.619		18.000		33.41
12	MET	C ·	67.438		17.710		17.211		32.40
13	MET	0	67.335		17.745		16.010		34.44
14	MET	СВ	65.376		17.042		18.345		34.94
15	MET	CG	65.321	•	18.129		19.414		36.89
16	MET.	SD	63.595		18.598		19.738		37.03
17	MET	CE	63.053		17.127		20.689		35.19
18	GLU	Ν	68.060		18.625	,	17.893		31.60
19	GLU	CA	68.666		19.756		17.226		34.56
20	GLU	С	69.903		19.379		16.393		37.49
21	GLU	0	70.082		19.836	. '	15.267		37.90
22	GLU	СВ	68.955		20.859		18.236		36.57
23	GLU	CG	68.694		22.256		17.685		54.02
24	GLU	CD	68.602	,	23.271		18.792		84.59
25	GLU	OE1	68.338	*	22.965		19.970		60.30
26	GLU	OE2	68.826		24.499		18.340		56.97
<b>27</b> ·	LYS	N	70.740		18.506		16.928		34.63
28	LYS	CA	71.925		18.073		16.173		36.36
29	LYS	C	71.504		17.225		14.995		35.30
30	LYS	0	72.071		17.271		13.926		33.46
31.	LYS	CB	72.858		17.280		17.069		44.20
32	LYS	CG	73.694		18.196		17.980	•	95.46
33	LYS	CD	74.837		17.496		18.729		100.00
34	LYS	CE	74.640		17.419		20.241		98.28
35	LYS	NZ	73.758	. •	16.304	•	20.653		100.00
_36	GLU	_N	70.463		16.441		_15.234_		-32-09-
37	GLU	CA	69.894		15.573		14.227		31.58
38	GLU	C	69.285	٠.	16.367		13.096		33.03

39	GLU	0	69.380	•	16.076		11.917		33.46
40	GLU	CB	68.841		14.653		14.863		33.60
41	GLU	CG	67.823		14.152		13.814		51.32
42	GLU	CD	66.936		13.044		14.309		61.90
43	GLU	OE1	66.302		13.085		15.370		49.63
44	GLU	OE2	66.918		12.042		13.457		46.46
45	PHE	N	68.648		17.422		13.475		31.95
46	PHE	CA	68.008		18.269		12.488		32.19
47	PHE	C	69.072		18.712		11.539		37.60
48	PHE	0	68.928		18.630		10.309		32.52
49	PHE	СВ	67.340		19.508	,	13.152		32.26
50	PHE	CG	66.508		20.348		12.196		30.98
51	PHE	CD1	65.161		20.064		11.967		29.63
52	PHE	CD2	67.094		21.402		11.499		29.81
53	PHE	CE1	64.398		20.834		11.096		31.86
54	PHE	CÉ2	66.354	:	22.185		10.621		32.68
55	PHE	CZ	65.004		21.896		10.423		34.10
	GLU	N N	70.164		19.160		12.179		36.63
56	_				19.627		11.440		36.44
57	GLU	CA	71.310				10.519		37.22
58	GLU	С	71.889		18.598		9.312 41	1 42	31.22
59	GLU	0	72.034		18.827			1.43	40.20
60	GLU	CB	72.309		20.346		12.308		40.39
61	GLU	CG	71.810		21.794		12.529		71.18
62	GLU	CD	71.946		22.266		13.953		100.00
63	GLU	OE1	72.735		21.752		14.751		100.00
64	GLU	OE2	71.139		23.288		14.223		100.00
65	GLN	N	72.140		17.413		11.024		31.25
66	GLN	CA	72.622		16.443		10.091		30.97
67	GLN	С	71.717		16.227			7.58	
68	GLN	0	72.187		16.205		7.798 35	5.23	
69	GLN	CB	72.828		15.118		10.746		32.09
70	GLN	CG	73.907		15.196		11.804		59.96
71	GLN	CD	74.286		13.786		12.123	,	100.00
72	GLN	OE1	73.653		12.854		11:579		100.00
73	GLN	NE2	75.309		13.631		12.975		100.00
74	- ILE	N	70.403		16.026		9.164 37	7.32	
75	ILE	CA	69.439		15.745		8.091 33	3.95	
76	ILE	C	69.451		16.857		7.112 35		
77	ILE	Ö	69.497		16.713			2.60	. *
78	ILE	СВ	68.007		15.516	•		3.88	
79	ILE	CG1			14.281		9.450 33		
80	ILE	CG2	67.062		15.263		7.427 27		
81	ILE	CD1	66.734		14.241		10.340		41.16
82	ASP	N	69.392		17.990		7.705 31	1 68	
	ASP	CA	69.374		19.138		6.893 34		
-83 - 24			70.643		19.138		6.028 45		
84	ASP	U	10.043		13.133		0.020 40	,.00	

85 87 88 90 91 93 94 95 97 98 99 101 103 104 105 107 108 109 111 113 114 115 116 117 118 119 119 119 119 119 119 119 119 119	ASPPSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSS	O C C C C C C C N N C C O C O N C C O C O	70.614 69.131 67.950 67.080 67.978 71.777 73.008 73.035 73.357 74.246 74.736 75.455 76.327 75.740 72.692 72.713 71.575 71.464 72.726 72.148 70.729 69.560 68.809 68.437 68.578 67.894 66.529 66.192 68.822 69.368 67.894 66.529 66.192 68.822 69.368 65.719 64.390 63.521 62.773 63.700 64.317 65.208 64.101 65.553 64.916 63.346 64.926	19.383 20.360 21.114 20.557 22.408 19.003 19.033 17.928 18.177 19.032 17.622 17.518 16.267 15.593 15.604 14.678 14.225 14.254 16.629 16.743 15.410 14.883 14.814 13.545 13.420 12.328 12.442 12.791 14.495 14.413 13.375 12.639 15.754 16.654 17.670 16.581 18.274 17.588 15.749 17.778	4.778 46.01 7.773 36.60 7.297 41.28 6.700 43.16 7.544 44.81 6.699 41.67 5.954 43.82 4.864 46.27 3.709 45.13 6.859 48.59 7.242 93.12 8.604 100.00 8.797 100.00 9.683 100.00 5.240 40.90 4.309 41.87 3.324 48.03 2.502 46.24 4.998 47.13 6.292 62.32 3.441 45.84 2.595 46.85 2.577 51.28 1.532 56.29 3.724 41.13 3.650 37.67 4.344 33.83 4.793 32.66 4.043 39.40 5.268 49.16 4.371 28.03 4.947 24.69 4.242 30.79 4.915 30.44 4.922 24.38 5.925 24.63 5.685 27.42 7.359 22.65 6.893 27.47 7.946 28.99 8.195 21.13 9.345 24.85	
			63.385	15.932	9.554 20.43	
127	TRP	CZ3				ne.
128	TRP	CH2	64.168	16.938	10.115 21.	UO
-129	ALA		63.620	13.268	2.876 26.09	
130	ALA	CA	62.799	12.286	2.153 24.22	

			· ·	•		
131	ALA	С	63.096	10.865	2.571	28.29
132	ALA	0	62.214	10.029	2.737	27.06
133	ALA	CB	62.920	12.477	0.652	25.69
		N	64.363	10.580		26.20
134	ALA					20.20
135	ALA	CA	64.704	9.238 3.195		
136	ALA	С	64.197	8.932 4.602		
137	ALA	0	63.581	7.885 4.927	27.14	
138	ALA	CB	66.210	9.022 3.107	26.43	*
139	ILE	N	64.482	9.876 5.467	28.04	
140	ILE	CA	64.042		25.50	
141	ILE	C·	62.562	9.449 6.863	27.81	
		Ö	62.053	8.525 7.520		,
142	ILE					28.83
143	ILE	CB	64.267	11.063		
144	ILE	CG1	65.751	11.246	7.430	30.75
145	ILE.	CG2	63.815	11.019	8.941	29.55
146	ILE	CD1	66.368	10.532	8.621	40.82
147	TYR	N	61.873	10.317	6.156	25.16
148	TYR	CA	60.436	10.229	6.111	24.00
149	TYR	С	59.987	8.882 5.562	28.75	
150	TYR	Ö	59.127	8.228 6.160	24.95	
151	TYR	СВ	59.814	11.445		23.47
152	TYR	CG	58.290	11.319		24.07
	TYR	CD1	57.449	11.372		25.26
153				11.154		24.94
154	TYR	CD2	57.674			
155	TYR	CE1	56.060	11.231		22.60
156	TYR	CE2	56.279	11.044		24.32
157	TYR	CZ	55.470	11.103		22.19
158	TYR	ОН	54.112	11.014		21.43
159	GLN	N	60.604	8.446 4.440	26.68	
160	GLN	CA	60.271	7.134 3.869	25.28	
161	GLN	C	60.553	6.006 4.861	26.17	
162	GLN	O :	59.857	4.992 4.963	26.54	• • •
163 ·	GLN	CB ·	61.021	6.871 2.543	27.78	
164	GLN	CG	62.409	6.217 2.796	84.03	el e
165	GLN	CD	63.607	6.501 1.839		
166	GLN	OE1	64.737	6.062 2.164		
167	GLN	NE2	63.414	7.188 0.676		•
				6.176 5.640		•
168	ASP	N	61.596			
169	ASP	CA	61.862	5.128 6.590		
170	ASP	C .	60.721	4.997 7.550		
171	ASP	0	60.290	3.884 7.886		
172	ASP	CB	63.220	5.284 7.314		
173	ASP	CG	64.331	5.565 6.310		·
174	ASP	OD1	64.144	5.579 5.099		
-175-	-ASP	-OD2-	65-510	5.815 6.842	-91:.88	
176	ILE	N	60.210	6.141 7.974	24.27	
-,	_	•	•	_		

177	ILE	CA	59.060	6.052 8.889 24.52	
178	ILE	C	57.903	5.367 8.255 24.70	
179	ILE	Ö	57.252	4.522 8.841 25.35	
180	ILE	СВ	58.619	7.415 9.401 27.95	
181	ILE	CG1	59.610	7.838 10.487	28.44
182	ILE	CG2	57.225	7.315 9.999 23.89	
183	ILE	CD1	59.930		27.02
184	ARG	N	57.646	5.725 7.020 22.44	
185	ARG	CA	56.511	5.098 6.330 22.70	
186	ARG	C	56.702	3.601 6.226 26.26	
187	ARG	Ŏ	55.761	2.788 6.333 23.08	
188	ARG	СВ	56.366	5.662 4.905 27.59	
189	ARG	CG	55.825	7.104 4.773 27.34	
190	ARG	CD	55.228	7.330 3.376 30.48	
191	ARG	NE	54.182	8.369 3.362 86.57	
192	ARG	CZ	53.614	8.942 2.268 100.00	)
193	ARG	NH1	53.954	8.615 1.006 100.00	
194	ARG	NH2	52.685	9.890 2.445 33.19	
195	HIS	N	57.967	3.235 5.974 26.18	
196	HIS	CA	58.297	1.840 5.840 28.26	
190	HIS	C	57.980	0.991 7.099 30.43	
		0	57.474	-0.1797.075 22.68	
198	HIS	CB	59.770	1.728 5.431 32.89	
199	HIS		60.149	0.296 5.206 42.37	
200	HIS	CG		-0.5046.250 47.99	
201	HIS	ND1	60.626	-0.4744.078 47.47	
202	HIS	CD2	60.082	-1.726 5.745 48.95	
203	HIS	CE1	60.816	-1.7474.449 48.75	
204	HIS	NE2	60.502	1.588 8.255 30.06	
205	GLU	N	58.321		
206	GLU	CA	58.143	0.866 9.524 28.09	27.30
207	GLU	C·	56.806	1.041 10.196	27.94
208	GLU	0	56.503	0.399 11.193	
209	GLU	CB	59.244	1.273 10.531	30.97
210	GLU	CG	60.629	1.547 9.904 54.48	100.00
211	GLU	CD	61.444	2.586 10.685	100.00
212	GLU	OE1	61.742	2.444 11.872	100.00
213	GLU	OE2	61.812	3.644 9.973 100.00	J
214	ALA	N	55.999	1.936 9.673 21.78	40.05
215	ALA	CA	54.703	2.217 10.276	19.05
216	ALA	C		0.959 10.372	26.56
217	ALA	0	53.939	0.125 9.462 25.40	
218	ALA	СВ	53.944	3.236 9.423 20.42	04:44
219	SER	N	53.081	0.847 11.465	21.44
220	SER	CA	52.234	-0.307 11.732	19.39
	-SER -		51.225	-0.517 10.663	27.89
222	SER	0	50.657	0.440 10.137	25.51
					9. 4

223	SER	CB	51.412	-0.049 12.974	21.80
224	SER	OG	52.257	0.317 14.021	26.89
225	ASP	N	50.935	-1.779 10.428	27.10
226	ASP	CA	49.936	-2.1299.448 29.07	
227	ASP	C	48.895	-2.997 10.125	30.02
	ASP	0	49.166	-4.13310.484	31.00
228				-2.786 8.250 33.50	31.00
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230	ASP	CG	49.690	-3.3487.216 50.19	
231	ASP	OD1	48.519	-3.0407.156 46.03	
232	ASP	OD2	50.278	-4.1856.378 67.71	00.70
233	PHE	N	47.737	-2.422 10.384	20.70
234	PHE	CA	46.675	-3.127 11.085	19.53
235	PHE	С	45.446	-3.117 10.216 <sup>-</sup>	25.93
236	PHE	0	45.307	-2.2819.357 28.17	
237	PHE	CB	46.339	-2.422 12.436	19.46
238	PHE	CG	47.428	-2.504 13.514	18.83
239	PHE	CD1	47.752	-3.720 14.138	19.35
240	PHE	CD2	48.062	-1.346 13.989	17.72
241	PHE	CE1	48.753	-3.782 15.118	19.95
242	PHE	CE2	49.088	-1.384 14.939	21.08
243	PHE	CZ	49.410	-2.61115.530	20.14
244	PRO	N	44.534	-4.03110.446	23.52
245	PRO	CA	43.331	-4.115 9.640 21.50	
246	PRO	C	42.303	-3.0019.968 23.90	• •
247	PRO	0	42.217	-2.497 11.117	22.13
		CB	42.675	-5.448 10.030	22.86
248	PRO		43.276	-5.845 11.381	29.00
249	PRO	CG		-5.147 11.450	24.62
250	PRO	CD	44.623		24.02
251	CYS	N	41.517	-2.717.8.941 19.44	
252	CYS	CA	40.442	-1.753 8.931 21.26	
253	CYS	C	39.268	-2.405 8.253 24.35	
254	CYS	0	38.706	-1.8907.289 23.90	
255	CYS	CB	40.832	-0.547 8.032 24.83	
256	CYS	SG	42.442	0.202 8.391 31.37	
257	ARG	N	38.910	-3.578 8.709 21.13	
258	ARG	CA	37.877	-4.2818.029 19.87	
259	ARG	С	36.558	-3.5828.150 24.13	
260	ARG	0	35.758	-3.5697.221 22.22	
261	ARG	CB	37.842	-5.7068.532 29.48	
262	ARG	CG	36.735	-5.804 9.576 76.36	
263	ARG	CD	36.827	-7.051 10.450	94.77
264	ARG	NE	36.033	-6.953 11.671	78.60
265	ARG	CZ	34.718	-6.843 11.637	91.25
266	ARG	NH1	34.073	-6.801 10.466	65.21
267	ARG	NH2	34.032	-6.768 12.785	88.97
268	VAL	N:	36.307	-2.9619.284 19.02	
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269 270 271 272 273 274 275 276 277 278 281 282 283 284 285 286 289 291 292 293 294 295 296 297 298 299 301 302 303 304 305 306 307 308 309 309 309 309 309 309 309 309 309 309	VAL VAL VAL VAL ALA ALA SSSSSSUU LEU LEU LEU PROOOOO PROPROSSSSSSSSSSSSSSSSSSSSSSSSS	CA COCGON CCOCGOCONN CCOCGON C	35.034 34.925 33.923 34.726 33.338 34.778 35.964 35.933 35.664 35.129 37.320 36.118 35.993 34.718 34.497 37.201 38.442 38.066 39.121 38.518 33.855 32.594 31.830 31.754 31.721 32.743 31.726 31.721 32.743 31.726 31.131 30.345 29.470 29.435 29.470 29.435 29.470 29.435 29.470 29.435 29.470 29.435 29.470 29.435 29.790 31.159 28.732 27.805 28.449 27.751 26.915 27.683 26.911	-2.288 9.372 17.46 -1.135 8.397 21.91 -0.950 7.725 25.41 -1.775 10.740 -1.205 10.712 -2.908 11.719 -0.367 8.277 17.06 0.744 7.364 17.17 0.295 5.949 26.79 1.038 5.135 23.44 1.378 7.356 16.97 -0.899 5.645 21.85 -1.403 4.299 23.03 -2.121 4.012 26.65 -2.565 2.898 29.21 -2.228 3.868 28.38 -1.359 3.651 30.93 -0.075 2.926 50.71 0.512 1.999 52.15 1.459 1.033 53.44 -2.250 4.983 26.04 -2.885 4.664 24.42 -2.075 3.603 30.28 -0.856 3.588 26.94 -2.820 5.907 25.29 -4.091 6.733 31.05 -5.118 6.282 30.64 -3.789 8.213 25.65 -2.743 2.705 33.23 -0.856 3.588 26.94 -2.820 5.907 25.29 -4.091 6.733 31.05 -5.118 6.282 30.64 -3.789 8.213 25.65 -2.743 2.705 33.23 -0.982 2.155 29.80 0.086 1.598 28.50 -3.139 1.155 35.94 -4.443 1.815 42.24 -4.194 2.448 36.58 -1.278 3.191 29.93 -0.291 3.727 29.87 1.003 4.192 32.18 2.019 4.352 29.62 -0.840 4.835 28.98 -1.496 5.963 32.04 -1.359 7.260 40.42	17.65 17.50 19.97 3 4 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
307	LYS	0	27.751	2.019 4.352 29.62	2
310	LYS	.CD	26.911	-1.3597.260 40.42	2
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				12.938	2.016	13.76
362	ARG	CG .	37.024			
363	ARG	CD	37.420	14.200	2.774	16.51
364	ARG	NE	36.224	14.791	3.392	18.16
365	ARG	CZ	35.306	15.542	2.703	26.85
					1.381	20.84
366	ARG	NH1	35.365	15.811		
367	ARG	NH2	34.234	16.012	3.341	16.64
368	TYR	N	38.164	9.354 1.679	17.86	
369	TYR	CA	39.233	8.872 0.832	15.41	
		C	39.411	7.359 0.980	22.83	
370	TYR					
371	TYR	O	39.443	6.781 2.075	16.42	
372	TYR	CB	40.562	9.498 1.157	14.29	•
373	TYR	CG	40.539	11.006	1,106	18.83
374	TYR	CD1	40.543	11.605	-0.152	18.23
		CD2	40.543	11.799	2.270	
375	TYR					
376	TYR	CE1	40.467	12.989	-0.282	
377	TYR	CE2	40.519	13.195	2.150	
378	TYR	CZ	40.508	13.767	0.872	18.16
379	TYR	ОН	40.491	15.147	0.711	18.89
380	ARG	N	39.483	6.735 -0.189		
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384	ARG	CB	39.306	4.858 -1.720	21.50	
385	ARG	CG	40.427	4.051 -2.346	62.10	•
386	ARG	CD	41.233	4.684 -3.494		
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387	ARG	NE	42.611			
388	ARG	CZ	43.771		3 100.00	J
389	ARG	NH1	43.842	6.149 -3.796		
390	ARG	NH2	44.910	4.155 -3.451	95.71	
391	ASP	N	41.862	5.576 0.492	14.95	
392	ASP	CA	43.082	5.093 1.065		
				5.554 2.490		
393	ASP	C	43.336			
394	ASP	0	44.434	5.386 3.007		
395	ASP	CB	44.260	5.583 0.229	16.63	
396	ASP	CG	44.232	7.082 0.082	22.24	
397	ASP	OD1	43.217	7.738 0.070	21.30	
•		OD2	45.394	7.561 -0.238		
398	ASP			6.156 3.118		
399	VAL	N	42.347			
400	VAL	CA	42.521	6.606 4.512		
401	VAL	С	41.410	6.066 5.346	14.75	
402	VAL	0	40.238	6.504 5.271	14.44	
403	VAL	СВ	42.451	8.123 4.686		
		CG1	42.721	8.517 6.182		
404	VAL					
405	-VAL			8.755 3.753		
406	SER	N:	41.767	5.113 6.158	15.99	

407 408 409 410	SER SER SER SER	CA C O CB	40.760 41.244 42.424 40.514	4.485 6.993 17.42 4.320 8.405 19.32 4.173 8.656 18.77 3.028 6.484 22.19	
411	SER	OG	40.054	3.029 5.131 21.85	
412	PRO	N CA	40.292 40.684	4.229 9.316 17.04 3.951 10.686	14.95
413 414	PRO PRO	CA	40.884	2.428 10.873	21.72
415	PRO	0	40.331	1.571 10.353	21.93
416	PRO	СВ	39.423	4.252 11.548	14.87
417	PRO	CG .	38.238	4.153 10.604	18.96
418	PRO	CD	38.800	4.338 9.177 15.06	
419	PHE	N	42.019	2.096 11.691	19.00
420	PHE	CA	42.266	0.711 12.046	16.26
421	PHE	C	41.099	0.239 12.907	20.51
422	PHE	0	40.517	0.996 13.712	18.21
423	PHE	CB (	43.484	0.629 12.972	16.62
424	PHE	CG	44.768	0.998 12.290	16.69
425	PHE	CD1	45.003	0.566 10.991	17.42
426	PHE	CD2	45.748	1.751 12.951	16.19
427	PHE	CE1	46.217	0.883 10.383	17.05
428	PHE	CE2	46.957	2.090 12.351	17.24
429	PHE	CZ	47.157	1.686 11.030	15.76
430	ASP	N	40.774	-1.056 12.819	19.69
431	ASP	CA	39.725	-1.541 13.645	19.38
432	ASP	C	40.050	-1.394 15.135	22.03 20.90
433	ASP	0	39.169	-1.155 15.966 2.022 12.221	21.07
434	ASP	CB	39.442	-3.033 13.331 -3.204 11.964	26.66
435	ASP	CG OD1	38.887 38.132	-2.404 11.443	25.86
436	ASP ASP	OD1 OD2	39.391	-4.217 11.336	29.95
437 438	HIS	N	41.288	-1.623 15.513	16.04
439	HIS	CA	41.509	-1.608 16.924	16.58
440	HIS	C .	41.355	-0.281 17.634	25.32
441	HIS	Ö	41.100	-0.239 18.870	24.53
442	HIS	CB .	42.856	-2.240 17.272	15.76
443	HIS	CG	44.037	-1.338 17.088	17.63
444	HIS	ND1	44.449	-0.441 18.079	19.17
445	HIS	CD2	44.890	-1.230 16.041	16.56
446	HIS	CE1	45.560	0.159 17.644	17.00
447	HIS	NE2	45.831	-0.298 16.427	17.69
448	SER	Ν	41.535	0.808 16.896	18.44
449	SER	CA	41.467	2.102 17.571	17.07
450	SER	C	40.307	2.966 17.101	24.02
	-SER-		-40.171	- 4.125-17.523	-19:51 17:01
452	SER	CB	42.776	2.867 17.350	17.91

453 454	SER ARG	OG N	43.130 39.469	2.848 15.967 2.429 16.223	17.63 18.61
455 456	ARG ARG	CA C	38.403 37.438	3.278 15.711 3.714 16.763	17.44 21.37
456 457	ARG	0	37.430	2.969 17.729	21.12
458	ARG	СВ	37.602	2.640 14.577	20.23
459	ARG	CG	36.621	1.515 15.009	21.54
460	ARG	CD	. 35.968	0.725 13.835	24.77
461	ARG	NE	34.948	-0.234 14.300	23.61
462	ARG	CZ	33.667	0.024 14.419	28.43
463	ARG	NH1	33.166	1.215 14.106	17.26
464	ARG	NH2	32.865	-0.945 14.886	22.24
465	ILE	Ν	36.814	4.891 16.529	20.42
466	ILE	CA	35.777	5.390 17.455	17.73
467	ILE	С	34.431	4.780 17.042	23.53
468	ILE	0	34.021	4.855 15.864	19.57
469	ILE	СВ	35.640	6.925 17.449	18.61
470	ILE	CG1	36.949	7.648 17.816	16.90
471	ILE	CG2	34.493	7.340 18.369	19.54
472	ILE	CD1	37.390	7.446 19.280	23.50
473	LYS	N	33.724	4.181 18.014	17.77 19.44
474	LYS	CA	32.479 31.329	3.661 17.638 4.585 18.080	23.14
475 476	LYS LYS	C O	31.329	5.073 19.222	23.14
476 477	LYS	CB	32.343	2.312 18.288	25.11
478	LYS	CG	33.271	1.269 17.706	28.50
479	LYS	CD	32.904	-0.078 18.301	42.01
480	LYS	CE	34.060	-1.057 18.404	50.09
481	LYS	NZ	33.628	-2.377 18.900	65.13
482	LEU	N	30.358	4.822 17.187	19.37
483	LEU	CÀ	29.173	5.650 17.536	19.21
484	LEU	<b>C</b> .	28.311	4.790 18.451	30.47
485	LEU	0	28.311	3.555 18.264	27.66
486	LEŲ	CB	28.346	5.917 16.283	18.17
487	LEU	CG	29.225	6.632 15.268	20.04
488	LEU	CD1	28.533	6.828 13.952	17.50
489	LEU	CD2	29.630	7.998 15.864	17.41
490	HIS	N	27.616	5.406 19.435	24.92
491	HIS	CA	26.790	4.616 20.333	23.96
492	HIS	C	25.439	4.506 19.717	34.63
493	HIS	0	24.491	5.195 20.064	38.67
494	HIS	CB	26.695	5.217 21.719	24.41
495 406	HIS HIS	CG ND1	28.030 28.197	5.381 22.372 6.121 23.570	29.98 33.91
496 497	HIS-	-CD2-	-29:258: - ·	4-908-21.994	- 31.10
49 <i>1</i> 498	HIS	CE1	29.504	6.052 23.881	33.12
430	1110		23.JU <del>4</del>	0.002 20.00 r	00.12

499	HIS	NE2	30.159	5.332 22.951 3.685 18.712	32.31 38.08
500	GLN	N	25.367 24.103	3.522 18.003	41.99
501	GLN.	CA C	24.103 \ 24.077	2.125 17.437	46.43
502 503	GLN GLN	0	2 <del>4</del> .077 25.111	1.523 17.202	42.03
503	GLN	CB	23.751	4 004 40 0E0	43.94
505	GLN	CG	24.597	4.425 15.691	50.12
506	GLN	CD	24.397	5.284 14.508	78.89
507	GLN	OE1	24.235	6.526 14.622	72.88
508	GLN	NE2	23.934	4.636 13.354	62.05
509	GLU	N	22.876	1.603 17.293	50.52
510	GLU	CA	22.676	0.234 16.832	53.51
511	GLU	C -	22.637	0.058 15.328	53.08
512	GLU	Ö	23.006	-0.999 14.825	48.73
513	GLU	СВ	21.441	-0.418 17.510	56.24
514	GLU	CG	21.550	-0.429 19.051	74.99
515	GLU	CD	20.383	-1.136 19.705	100.00
516	GLU	OE1	19.203	-0.939 19.368	100.00
517	GLU	OE2	20.768	-2.009 20.636	100.00
518	ASP	N	22.170	1.083 14.619	50.94
519	ASP	CA	22.101	0.990 13.183	52.56
520	ASP	C	23.492	0.706 12.532	50.81
521	ASP	0	23.723	-0.285 11.795	54.32
522	ASP	CB	21.388	2.250 12.634	57.77
523	ASP	CG.	21.808	2.668 11.243	94.02
524	ASP	OD1	21.577	1.977 10.250	99.87
525	ASP	OD2	22.439	3.847 11.214	100.00
526	ASN	N	24.444	1.597 12.808	34.60
527	ASN	CA	25.773	1.475 12.236	28.07
528	ASN	С	26.669	2.345 13.088	29.15
529	ASN	0	26.536	3.556 13.101	30.48
530	ASN	CB	25.734	2.022 10.803	19.62
531	ASN	CG	27.024	1.823 10.062	27.76
532	ASN	OD1	28.067	1.547 10.679	22.79
533	ASN	ND2	26.967	1.995 8.729 21.65	
534	ASP	N .	27.600	1.752 13.806	22.25
535	ASP	CA	28.430	2.534 14.667	20.02
536	ASP	С	29.676	3.124 14.011	22.15
537	ASP	0	30.575	3.603 14.710	22.95
538	ASP	CB	28.858	1.548 15.757	21.84
539	ASP	CG	29.803	0.461 15.282	26.22
540	ASP	OD1	30.328	0.421 14.195	26.63
541	ASP	OD2	30.146	-0.355 16.235	35.64
542	TYR	Ν	29.794	3.001 12.697	19.04
543	TYR	CA	31.033	3.440 12.034	16.92
544	TYR	С	31.184	4.931 11.701	22.03

		_		5.004.44.407	00.04	
545	TYR	0	30.325	5.601 11.107	22.84	
546	TYR	CB		2.733 10.691	16.06	
547	TYR			3.169 9.987 17.61	20.91	
548	TYR				20.91	
549	TYR	CD2	32.403	4.043 8.886 16.20		
550	TYR	CE1	34.857	3.065 9.828 19.50		
551	TYR	CE2	33.596	4.460 8.282 13.17 3.973 8.769 14.97		
552	TYR	CZ	34.818	4.360 8.214 18.92		
553	TYR	OH		5.440 12.061	19.32	
554 555	ile	N	32.319	6.780 11.682	17.75	
555	ILE	CA	32.752	6.676 11.366	16.65	
556	ILE	C			14.78	
557	ILE	0		5.912 11.984	18.72	
558	ILE	CB		7.883 12.709		•
559	ILE			9.269 12.196	17.13	
560	ILE		33.132			16.24
561	ILE			10.364 13.106		16.24
562	ASN	N			14.05	
563	ASN	CA		7.408 10.072	12.04	
564	ASN	C	,	8.263 11.062	18.49	
565		,0	37.028	9.462 10.808		
566	ASN		36.392	7.908 8.637 10.31		
567			37.806	7.615 8.161 18.51		
568	ASN		38.803	7.928 8.840 17.25		
569	ASN	ND2	-	7.096 6.948 13.86		÷
570	ALA	Ν	37.206	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	15.81	
571	ALA	CA	37.822	8.363 13.318	15.05	
572	ALA	С	38.551		17.50	
573	ALA		38.093	6.297 14.414	15.99	
574	ALA		36.768	9.111 14.121	15.21	
575	SER	N	39.670		13.78	
576	SER	CA	40.518	7.002 15.565	14.32	
577	SER	С	40.915	7.650 16.857	16.94	
578	SER	0	41.210	8.834 16.908	18.12	
579	SER	CB	41.859	6.766 14.820	15.23	
580	SER	OG	41.642	6.232 13.488	17.37	
581	LEU	N	41.046	6.853 17.898	15.51	
582	LEU	CA	41.503	7.396 19.161	15.13	
583	LEU	С	43.015	7.175 19.229	22.22	
584	LEU	0	43.454	6.029 19.128	21.68	•
585	LEU	CB	40.801	6.641 20.374	16.59	
586	LEU	CG ·	41.333	6.988 21.784	19.94	
587	LEU	CD1	41.053	8.438 22.118	20.86	
588	LEU	CD2	40.611	6.204 22.847	21.37	
589	ILE	N	43.797	8.247 19.421	17.07	
590	ILE	CA	45.219	8.168 19.506	16.19	
				7	. * *	

591	ILE	C ·	45.524	8.335 20.99	5	24.53	
592	ILE	Ö	45.338	9.380 21.56		22.94	
		_		9.330 18.79		18.65	
593	ILE	CB	45.845				
594	ILE	CG1	45.927	9.229 17.28		18.37	
595		CG2	47.265	9.378 19.29		21.51	i
596	ILE	CD1	44.791	8.564 16.61	1 .	25.42	
597	LYS	N	45.955	7.285 21.66	4	20.46	
598	LYS	CA	46.162	7.350 23.09	2	20.79	
599	LYS	C	47.630	7.299 23.39		24.25	
600	LYS	0	48.236	6.260 23.15		24.06	•
		CB	45.396	6.160 23.69		24.07	
601	LYS						•
602	LYS	CG	44.960	6.286 25.15		48.81	
603	LYS	CD	44.128	5.081 25.61		70.96	
604	LYS	CE	44.276	4.756 27.10		99.34	
605	LYS	NZ	44.076	3.328 27.44	5	100.00	
606	MET	Ν	48.201	8.455 23.82	2	22.51	
607	MET	CA	49.625	8.581 24.12	4	20.07	
608	MET	C	49.859	8.290 25.59		27.14	
609	MET.	0	49.758	9.141 26.46		24.51	
				9.882 23.64		19.62	
610	MET	CB	50.266				14 42
611	MET	CG	50.032	10.097	22.162		21.13
612	MET	SD	50.570	8.761 21.08		23.27	
613	MET	CE	52.316	9.093 21.05		18.81	
614	GLU	N ,	50.136	7.023 25.83	0	28.15	
615	GLU	CA	50.280	6.525 27.16	0	29.78	`
616	GLU	С	51.248	7.321 28.03	0	34.27	
617	GLU	0	50.881	7.991 29.01		33.62	•
618	GLU	CB	50.621	5.054 27.05		30.77	
619	GLU	CG	50.491	4.307 28.37		42.42	
						86.17	
620	GLU	CD	50.541	2.833 28.16			
621	GLU	OE1	51.464	2.282 27.58		100.00	
622	GLU	OE2	49.454	2.226 28.58		100.00	
623	GLU	N	52.506	7.246 27.64		30.68	
624	GLU	CA	53.546	7.943 28.39	6	31.16	
625	GLU	С	53.243	9.397 28.51	8	35.90	
626	GLU	0	53.388	9.913 29.56	7	36.43	
627	GLU	ĊВ	54.865	7.737 27.68	1	33.02	
628	GLU	CG	56.142	8.220 28.38		46.68	
629	GLU		57.242	8.086 27.35		78.67	
630	GLU	OE1	57.023	7.823 26.16		59.52	
631	GLU	OE2	58.437	8.258 27.83		84.75	
632	ALA	N	52.800	10.068	27.441		34.48
633	ALA	CA	52.488	11.493	27.524		31.47
634	ALA	С	51.242	11.736	28.308		33.12
635	ALA	0	51.026	12.820	28.781	3	31.13
636	ALA	CB	52.294	12.082	26.132	3	31.22
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GLN	N	50.354	10.764	28.383	32.47
		49.116	11.014	29.114	35.37
		48.196	12.076	28.454	39.78
			13.026	29.072	40.91
			11.386	30.560	39.09
			10.260	31.290	72.08
				7 100.00	0
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				,	
		47.979	11.913	27.140	28.50
		47.086	12.751	26.374	23.58
					24.24
			·		26.56
		,		25.722	23.27
			,		26.99
					26.70
					23.46
					24.69
					22.83
					23.63
					19.60
					18.87
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					20.11
					21.80
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					21.22
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					21.01
				19.695	20.98
				19.707	18.74
				20.298	16.73
					17.39
				20.511	16.87
				21.079	18.92
					17.15
					23.78
					23.03
			· ·		17.07
				•	16.38
					19.66
					19.42
					21.59
			at the second se		21.73
					19.23
				20.239	19.35
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	NNNNNNNNGGGGGGGGGGGKRRRRRRRRRRRRRRRRRRR	GLN CA CB GLN CB CD	GLN CA 49.116 GLN C 48.196 GLN O 47.700 GLN CB 49.434 GLN CG 50.174 GLN CD 49.157 GLN OE1 48.700 GLN NE2 48.738 ARG N 47.979 ARG CA 47.086 ARG C 46.524 ARG O 47.232 ARG CB 47.779 ARG CG 46.780 ARG CD 47.361 ARG NE 47.293 ARG CZ 47.954 ARG NH1 48.785 ARG NH2 47.826 SER N 45.249 SER CA 44.632 SER C 44.089 SER CA 44.632 SER C 44.089 SER O 43.869 SER CB 43.455 SER OG 43.930 TYR N 43.820 TYR CA 43.211 TYR C 42.381 TYR C 42.381 TYR C 42.381 TYR C 42.381 TYR CB 44.261 TYR CG 45.522 TYR CD1 46.618 TYR CB 44.261 TYR CB 44.261 TYR CG 45.522 TYR CD1 46.618 TYR CD2 45.619 TYR CB 44.261 TYR CB 44.261 TYR CB 44.261 TYR CB 45.522 TYR CD1 46.618 TYR CD2 45.619 TYR CB 47.790 TYR CE2 46.773 TYR CZ 47.854 TYR OH 49.001 ILE N 41.328 ILE CA 40.554 ILE CB 39.027 ILE CB 39.027 ILE CB 39.027 ILE CB 39.027 ILE CB 38.653 ILE CG2 38.255	GLN CA 49.116 11.014 GLN C 48.196 12.076 GLN O 47.700 13.026 GLN CB 49.434 11.386 GLN CG 50.174 10.260 GLN CD 49.157 9.374 31.95 GLN OE1 48.700 9.687 33.09 GLN NE2 48.738 8.341 31.20 ARG N 47.979 11.913 ARG CA 47.086 12.751 ARG C 46.524 11.885 ARG O 47.232 11.052 ARG CB 47.779 13.904 ARG CG 46.780 14.899 ARG CD 47.361 16.299 ARG NE 47.293 17.043 ARG CZ 47.954 18.135 ARG NH1 48.785 18.663 ARG NH2 47.826 18.721 SER N 45.249 12.005 SER CA 44.632 11.302 SER C 44.089 12.361 SER O 43.869 13.515 SER CB 43.455 10.478 SER OG 43.930 9.552 25.250 TYR N 43.820 11.959 TYR CA 43.211 12.812 TYR C 42.381 11.892 TYR	GLN CA 49.116 GLN C 48.196 GLN C 48.196 GLN C 48.196 GLN CB 49.434 GLN CG 50.174 GLN CD 49.157 GLN CD 49.157 GLN OE1 48.700 GLN NE2 48.738 ARG N 47.979 ARG CA 47.086 ARG CB 47.779 ARG CB 47.786 ARG CB 47.779 ARG CB 47.861 ARG

683	LEU	Ν	41.336	11.645	16.006	15.32
684	LEU	CA	41.675	12.234	14.715	14.26
685	LEU	C	40.656	11.756	13.748	18.37
					13.738	16.49
686	LEU	0	40.389	10.571		
687	LEU	CB	43.066	11.848	14.198	13.33
688	LEU	CG	44.175	12.803	14.642	17.41
689	LEU	CD1	45.534	12.403	13.995	16.26
690	LEU	CD2	44.281	12.897	16.182	15.86
691	THR	N	40.050	.12.645	12.963	14.18
692	THR	CA	39.024	12.145	12.076	13.33
			39.116	12.910	10.755	16.62
693	THR	С				15.02
694	THR	0	39.788	13.946	10.696	
695	THR	CB	37.644	12.338	12.815	15.01
696	THR	OG1	36.550	11.816	12.090	15.33
697	THR	CG2	37.396	13.829	13.035	11.94
698	GLN	Ν	38.432	12.417	9.692 15.62	
699	GLN	CA	38.460	13.162	8.437 14.08	
700	GLN	C	37.474	14.358	8.490 16.44	
701	GLN	Ö	36.541	14.444	9.358 16.22	
701	GLN	СВ	38.017	12.224	7.323 13.86	
					7.547 18.12	,
703	GLN	CG -	36.562	11.804		
704	GLN	CD	35.915	10.847	6.544 19.07	
705	GLN	OE1	34.655	10.690	6.513 19.98	
706	GLN	NE2	36.756	10.142	5.818 11.61	
707	GLY	Ν	37.556	15.222	7.487 15.42	
708	GLY	CA	36.598	16.340	7.384 12.58	
709	GLY	C	35.252	15:724	7.143 16.22	
710	GLY	Ö	35.067	14.965	6.204 14.03	
711	PRO	N	34.274	16.005	7.986 16.60	
712	PRO	CA	32.950	15.389	7.745 15.78	
				15.529	6.317 20.52	
713	PRO	C	32.405			
714	PRO	0	32.677	16.525	5.642 19.37	
715	PRO	CB	32.010	16.071	8.734 15.66	
716	PRO	CG	32.902	16.749	9.800 17.67	
717	PRO	CD	34.316	16.772	9.281 14.44	
718	LEU	·N	31.620	14.505	5.898 16.55	
719	LEU	CA	30.962	14.416	4.631 14.43	
720	LEU	C	29.522	14.909	4.834 22.58	
721 .	LEU	Ö	29.029	14.985	5.933 20.15	
		CB	30.952	12.997	4.038 14.77	
722	LEU				3.728 18.92	
723	LEU	CG	32.352	12.481		
724	LEU	CD1	32.333	10.957	3.798 20.45	
725	LEU	CD2	32.799	12.968	2.329 19.10	
726	PRO	N	28.852	15.291	3.742 23.61	
727	PRO	CA	27.526		3.867- 23.49	
728	PRO	C	26.616	14.852	4.520 26.41	

729 730 731 732 733 734 735 736 737 738 739	PRO PRO PRO ASN ASN ASN ASN ASN ASN	O CB CD N CA C O CB CG OD1	25.574 27.017 28.057 29.290 26.973 26.068 26.608 25.978 25.861 27.174 28.160	15.258 16.030 15.523 15.276 13.586 12.632 12.193 11.361 11.386 10.691 11.365	4.934 30.17 2.454 25.04 1.480 29.04 2.312 23.33 4.607 21.62 5.237 17.84 6.557 22.95 7.233 23.70 4.362 18.60 3.998 24.49 3.684 24.35	
740	ASN	ND2	27.214	9.338 4.017 12.714	24.03 6.939 16.40	
741 742	THR THR	N CA	27.791 28.325	12.714	8.220 15.19	
743	THR	C	28.433	13.364	9.223 21.17	•
744	THR	0	29.095	13.234	10.219	19.70
745	THR	CB	29.694	11.529	8.112 18.74	•
746	THR	OG1	30.690	12.447	7.709 19.88	
747	THR	CG2	29.683	10.379	7.103 16.93	
748	CYS	N	27.783	14.491	8.970 16.82	
749	CYS	CA	27.883	15.573	9.943 18.20	
750	CYS	С	27.174	15.247	11.228	19.02
751	CYS	0	27.613	15.697	12.308	20.16
752	CYS	CB	27.325	16.925	9.410 19.21	
753	CYS	SG	28.252	17.458	7.951 23.17	17.50
754 755	GLY	N	26.054	14.501	11.125 12.370	17.59 16.40
755 756	GLY GLY	CA C	25.352 26.210	14.185 13.222	13.189	18.70
756 757	GLY	0	26.279	13.293	14.394	18.52
758	HIS	N	26.865	12.306	12.499	17.15
759	HIS	CA	27.754	11.319	13.158	17.16
760	HIS	C	28.925	12.017	13.833	20.07
761	HIS	Ö	29.404		14.889	20.41
762	HIS	СВ	28.393	10.342	12.120	18.16
763	HIS	CG	27.384	9.635 11.299	9 19.12	
764	HIS	ND1	27.704	9.228 9.989	22.15	
765	HIS	CD2	26.096	9.300 11.596	5 17.48	
766	HIS	CE1	26.596	8.639 9.509		
767	HIS	NE2	25.620	8.661 10.438		
768	PHE	N	29.491	13.003	13.166	17.15
769	PHE	CA	30.592	13.729	13.736	15.25
770	PHE	C	30.214	14.331	15.139	22.13
771	PHE	O	30.894	14.098	16.171	21.51 15.12
772	PHE	CB	31.025	14.822	12.736 13.307	-14.74
773 774	PHE	CG CD1	32.096 31.746	15.740 16.887	14.020	15.97
114	LLIE	ועט	J 1.740	10.007	17.020	10.01

777 PHE CE2 34.475 16.298 13.663 17.69 778 PHE CZ 34.095 17.428 14.398 14.91 779 TRP N 29.096 15.088 15.168 19.55 780 TRP CA 28.632 15.704 16.421 18.79 781 TRP C 28.147 14.656 17.418 21.85 782 TRP O 28.289 14.839 18.633 21.13 783 TRP CB 27.576 16.758 16.169 16.67 784 TRP CG 28.210 17.904 15.491 16.11 785 TRP CD1 27.962 18.318 14.214 18.99 786 TRP CD2 29.206 18.776 16.023 14.97 787 TRP NE1 28.768 19.412 13.916 18.96 788 TRP CE2 29.547 19.692 15.015 18.62 789 TRP CE3 29.867 18.880 17.250 16.41 790 TRP CZ3 30.823 19.869 17.428 16.16 792 TRP CH2 31.165 20.737 16.412 16.86 793 GLU N 27.626 13.536 16.911 17.79 794 GLU CA 27.268 12.489 17.811 16.48 795 GLU C 28.499 12.004 18.561 20.28 796 GLU C 28.499 12.004 18.561 20.28 797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.414 12.466 19.451 21.03 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CG 31.905 11.171 17.275 15.36 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 812 VAL C 31.027 14.648 21.239 20.80 813 VAL O 31.512 14.740 22.361 21.08 813 VAL C 31.027 14.648 21.239 20.81 814 VAL CB 31.555 61.117 19.308 18.34 815 VAL CG 32.560 16.297 18.112 16.59 810 VAL CB 31.555 16.117 19.308 18.34 815 VAL CB 31.576 16.117 19.308 18.34 815 VAL CG 32.560 16.297 18.112 16.59 810 VAL CB 32.560 16.297 18.112 16.59	775 776	PHE PHE	CD2 CE1	33.466 32.734	15.466 17.739	13.140 14.539	15.95 16.67
779 TRP N 29.096 15.088 15.168 19.55 780 TRP CA 28.632 15.704 16.421 18.79 781 TRP C 28.147 14.656 17.418 21.85 782 TRP O 28.289 14.839 18.633 21.13 783 TRP CB 27.576 16.758 16.169 16.67 784 TRP CG 28.210 17.904 15.491 16.11 785 TRP CD1 27.962 18.318 14.214 18.99 786 TRP CD2 29.206 18.776 16.023 14.97 787 TRP NE1 28.768 19.412 13.916 18.96 788 TRP CE2 29.547 19.692 15.015 18.62 789 TRP CE3 29.867 18.880 17.250 16.41 790 TRP CZ3 30.506 20.686 15.213 17.94 791 TRP CZ3 30.823 19.869 17.428 16.16 792 TRP CH2 31.165 20.737 16.412 16.86 793 GLU N 27.626 13.536 16.911 17.79 794 GLU CA 27.268 12.489 17.811 16.48 795 GLU C 28.499 12.004 18.561 20.28 796 GLU O 28.480 11.817 19.780 18.97 797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.916 12.180 20.552 17.41 806 MET CB 31.905 11.171 17.275 15.36 807 MET CG 33.296 10.921 17.815 15.60 808 MET CG 33.296 10.921 17.815 15.60 808 MET CG 33.296 10.921 17.815 15.36 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 812 VAL C 31.027 14.648 21.239 20.81 813 VAL O 31.555 16.117 19.308 18.34 815 VAL CG1 31.720 17.289 20.311 17.11 816 VAL CG2 32.560 16.297 18.112 16.59 819 TRP C 28.223 13.143 23.061 25.99							
780         TRP         CA         28.632         15.704         16.421         18.79           781         TRP         C         28.147         14.656         17.418         21.85           782         TRP         O         28.289         14.839         18.633         21.13           783         TRP         CB         27.576         16.758         16.169         16.67           784         TRP         CB         27.576         16.758         16.169         16.67           784         TRP         CD         28.210         17.904         15.491         16.11           785         TRP         CD1         27.962         18.318         14.214         18.99           786         TRP         CD2         29.206         18.776         16.023         14.97           787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         <							
781         TRP         C         28.147         14.656         17.418         21.85           782         TRP         O         28.289         14.839         18.633         21.13           783         TRP         CB         27.576         16.758         16.169         16.67           784         TRP         CG         28.210         17.904         15.491         16.11           785         TRP         CD1         27.962         18.318         14.214         18.99           786         TRP         CD2         29.206         18.776         16.023         14.97           787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE2         30.506         20.686         15.213         17.94           790         TRP         CZ3         30.823         19.869         17.428         16.16           791         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         <							
782         TRP O         28.289         14.839         18.633         21.13           783         TRP CB         27.576         16.758         16.169         16.67           784         TRP CG         28.210         17.904         15.491         16.11           785         TRP CD1         27.962         18.318         14.214         18.99           786         TRP CD2         29.206         18.776         16.023         14.97           787         TRP NE1         28.768         19.412         13.916         18.96           788         TRP CE2         29.547         19.692         15.015         18.62           789         TRP CE3         29.867         18.880         17.250         16.41           790         TRP CZ3         30.506         20.686         15.213         17.94           791         TRP CZ3         30.823         19.869         17.428         16.16           792         TRP CH2         31.165         20.737         16.412         16.86           793         GLU CA         27.268         12.489         17.811         16.48           795         GLU CB         26.630         11.293         17.103							
783 TRP CB 27.576 16.758 16.169 16.67 784 TRP CG 28.210 17.904 15.491 16.11 785 TRP CD1 27.962 18.318 14.214 18.99 786 TRP CD2 29.206 18.776 16.023 14.97 787 TRP NE1 28.768 19.412 13.916 18.96 788 TRP CE2 29.547 19.692 15.015 18.62 789 TRP CE3 29.867 18.880 17.250 16.41 790 TRP CZ3 30.506 20.686 15.213 17.94 791 TRP CZ3 30.823 19.869 17.428 16.16 792 TRP CH2 31.165 20.737 16.412 16.86 793 GLU N 27.626 13.536 16.911 17.79 794 GLU CA 27.268 12.489 17.811 16.48 795 GLU C 28.499 12.004 18.561 20.28 796 GLU O 28.480 11.817 19.780 18.97 797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CG 26.576 10.108 18.107 18.51 799 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.414 12.466 19.451 21.03 805 MET O 31.916 12.180 20.552 17.41 806 MET C 31.414 12.466 19.451 21.03 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 814 VAL CB 31.555 16.117 19.308 18.34 815 VAL CG 31.527 14.648 21.239 20.80 814 VAL CB 31.555 16.117 19.308 18.34 815 VAL CG 31.555 16.117 19.308 18.34 816 VAL CG 23.560 16.297 18.112 16.59 817 TRP N 29.760 14.465 21.061 23.24 818 TRP CA 28.891 14.345 22.223 25.45 819 TRP CC 29.223 13.4143 22.205 25.99							
784         TRP         CG         28.210         17.904         15.491         16.11           785         TRP         CD1         27.962         18.318         14.214         18.99           786         TRP         CD2         29.206         18.776         16.023         14.97           787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.69           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499			_				
785         TRP         CD1         27.962         18.318         14.214         18.99           786         TRP         CD2         29.206         18.776         16.023         14.97           787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CE2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CB         26.630							
786         TRP         CD2         29.206         18.776         16.023         14.97           787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CB         26.630         11.293         17.103         17.70           799         GLU         CD         25.985							
787         TRP         NE1         28.768         19.412         13.916         18.96           788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CB         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         <							
788         TRP         CE2         29.547         19.692         15.015         18.62           789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         C         28.499         12.004         18.561         20.28           797         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CG         26.576         10.108         18.107         18.51           800         GLU         OE1         25.635 <t< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td></t<>							
789         TRP         CE3         29.867         18.880         17.250         16.41           790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         C         28.499         12.004         18.561         20.28           796         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CG         26.576         10.108         18.107         18.51           799         GLU         OE1         25.635         8.939 16.304         24.52           801         GLU         OE2         25.828         7.845 18.231 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>							
790         TRP         CZ2         30.506         20.686         15.213         17.94           791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CB         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11							
791         TRP         CZ3         30.823         19.869         17.428         16.16           792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         C         28.480         11.817         19.780         18.97           797         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CG         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11.8							
792         TRP         CH2         31.165         20.737         16.412         16.86           793         GLU         N         27.626         13.536         16.911         17.79           794         GLU         CA         27.268         12.489         17.811         16.48           795         GLU         C         28.499         12.004         18.561         20.28           796         GLU         C         28.480         11.817         19.780         18.97           797         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CG         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11.855         17.813         16.81           803         MET         CA         30.873         11.42			CZ3		19.869	17.428	16.16
794 GLU CA 27.268 12.489 17.811 16.48 795 GLU C 28.499 12.004 18.561 20.28 796 GLU O 28.480 11.817 19.780 18.97 797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.414 12.466 19.451 21.03 805 MET O 31.916 12.180 20.552 17.41 806 MET CB 31.905 11.171 17.275 15.36 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 812 VAL C 31.027 14.648 21.239 20.80 813 VAL O 31.512 14.740 22.361 21.08 814 VAL CB 31.555 16.117 19.308 18.34 815 VAL CG1 31.720 17.289 20.311 17.11 816 VAL CG2 32.560 16.297 18.112 16.59 817 TRP N 29.760 14.465 21.061 23.24 818 TRP CA 28.891 14.345 22.223 25.45 819 TRP C 29.223 13.143 23.061		TRP	CH2	31.165	20.737	16.412	16.86
795 GLU C 28.499 12.004 18.561 20.28 796 GLU O 28.480 11.817 19.780 18.97 797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CG 26.576 10.108 18.107 18.51 799 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.414 12.466 19.451 21.03 805 MET O 31.916 12.180 20.552 17.41 806 MET CB 31.905 11.171 17.275 15.36 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 812 VAL C 31.027 14.648 21.239 20.80 813 VAL O 31.512 14.740 22.361 21.08 814 VAL CB 31.555 16.117 19.308 18.34 815 VAL CG1 31.720 17.289 20.311 17.11 816 VAL CG2 32.560 16.297 18.112 16.59 817 TRP N 29.760 14.465 21.061 23.24 818 TRP CA 28.891 14.345 22.223 25.45 819 TRP C 29.223 13.143 23.061 25.99		GLU	N	27.626	13.536	16.911	17.79
796         GLU         O         28.480         11.817         19.780         18.97           797         GLU         CB         26.630         11.293         17.103         17.70           798         GLU         CG         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11.855         17.813         16.81           803         MET         CA         30.873         11.423         18.408         15.42           804         MET         CA         31.916         12.180         20.552         17.41           805         MET         O         31.916         12.180         20.552         17.41           806         MET         CB         31.905         11.171         17.275         15.36           807         MET         CB         34.636         12.33	794	GLU	CA	27.268	12.489	17.811	16.48
797 GLU CB 26.630 11.293 17.103 17.70 798 GLU CG 26.576 10.108 18.107 18.51 799 GLU CD 25.985 8.904 17.457 36.95 800 GLU OE1 25.635 8.939 16.304 24.52 801 GLU OE2 25.828 7.845 18.231 22.85 802 MET N 29.622 11.855 17.813 16.81 803 MET CA 30.873 11.423 18.408 15.42 804 MET C 31.414 12.466 19.451 21.03 805 MET O 31.916 12.180 20.552 17.41 806 MET CB 31.905 11.171 17.275 15.36 807 MET CG 33.296 10.921 17.815 15.60 808 MET SD 34.527 10.644 16.486 19.45 809 MET CE 34.636 12.330 15.779 16.29 810 VAL N 31.345 13.741 19.059 17.81 811 VAL CA 31.809 14.761 19.961 15.34 812 VAL C 31.027 14.648 21.239 20.80 813 VAL O 31.512 14.740 22.361 21.08 814 VAL CB 31.555 16.117 19.308 18.34 815 VAL CG1 31.720 17.289 20.311 17.11 816 VAL CG2 32.560 16.297 18.112 16.59 817 TRP N 29.760 14.465 21.061 23.24 818 TRP CA 28.891 14.345 22.223 25.45 819 TRP C 29.223 13.143 23.061 25.99	795	GLU	С	28.499	12.004	18.561	20.28
798         GLU         CG         26.576         10.108         18.107         18.51           799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11.855         17.813         16.81           803         MET         CA         30.873         11.423         18.408         15.42           804         MET         C         31.414         12.466         19.451         21.03           805         MET         O         31.916         12.180         20.552         17.41           806         MET         CB         31.905         11.171         17.275         15.36           807         MET         CG         33.296         10.921         17.815         15.60           808         MET         SD         34.527         10.644         16.486         19.45           809         MET         CE         34.636         12.33	796	GLU	0	28.480	11.817	19.780	18.97
799         GLU         CD         25.985         8.904         17.457         36.95           800         GLU         OE1         25.635         8.939         16.304         24.52           801         GLU         OE2         25.828         7.845         18.231         22.85           802         MET         N         29.622         11.855         17.813         16.81           803         MET         CA         30.873         11.423         18.408         15.42           804         MET         CA         30.873         11.423         18.408         15.42           804         MET         C         31.414         12.466         19.451         21.03           805         MET         O         31.916         12.180         20.552         17.41           806         MET         CB         31.905         11.171         17.275         15.36           807         MET         CG         33.296         10.921         17.815         15.60           808         MET         SD         34.527         10.644         16.486         19.45           809         MET         CE         34.636         12.33	797	GLU	CB	26.630			
800       GLU       OE1       25.635       8.939       16.304       24.52         801       GLU       OE2       25.828       7.845       18.231       22.85         802       MET       N       29.622       11.855       17.813       16.81         803       MET       CA       30.873       11.423       18.408       15.42         804       MET       C       31.414       12.466       19.451       21.03         805       MET       O       31.916       12.180       20.552       17.41         806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.555 <t< td=""><td>798</td><td>GLU</td><td></td><td></td><td></td><td></td><td>18.51</td></t<>	798	GLU					18.51
801       GLU       OE2       25.828       7.845 18.231       22.85         802       MET       N       29.622       11.855       17.813       16.81         803       MET       CA       30.873       11.423       18.408       15.42         804       MET       C       31.414       12.466       19.451       21.03         805       MET       O       31.916       12.180       20.552       17.41         806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117							
802       MET       N       29.622       11.855       17.813       16.81         803       MET       CA       30.873       11.423       18.408       15.42         804       MET       C       31.414       12.466       19.451       21.03         805       MET       O       31.916       12.180       20.552       17.41         806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>							
803       MET       CA       30.873       11.423       18.408       15.42         804       MET       C       31.414       12.466       19.451       21.03         805       MET       O       31.916       12.180       20.552       17.41         806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       <							
804       MET       C       31.414       12.466       19.451       21.03         805       MET       O       31.916       12.180       20.552       17.41         806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560							
805         MET         O         31.916         12.180         20.552         17.41           806         MET         CB         31.905         11.171         17.275         15.36           807         MET         CG         33.296         10.921         17.815         15.60           808         MET         SD         34.527         10.644         16.486         19.45           809         MET         CE         34.636         12.330         15.779         16.29           810         VAL         N         31.345         13.741         19.059         17.81           811         VAL         CA         31.809         14.761         19.961         15.34           812         VAL         C         31.027         14.648         21.239         20.80           813         VAL         O         31.512         14.740         22.361         21.08           814         VAL         CB         31.555         16.117         19.308         18.34           815         VAL         CG1         31.720         17.289         20.311         17.11           816         VAL         CG2         32.560         16.							
806       MET       CB       31.905       11.171       17.275       15.36         807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891							
807       MET       CG       33.296       10.921       17.815       15.60         808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223							
808       MET       SD       34.527       10.644       16.486       19.45         809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       -TRP       C       29.223       13.143       23.061       25.99							
809       MET       CE       34.636       12.330       15.779       16.29         810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
810       VAL       N       31.345       13.741       19.059       17.81         811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
811       VAL       CA       31.809       14.761       19.961       15.34         812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
812       VAL       C       31.027       14.648       21.239       20.80         813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
813       VAL       O       31.512       14.740       22.361       21.08         814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
814       VAL       CB       31.555       16.117       19.308       18.34         815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
815       VAL       CG1       31.720       17.289       20.311       17.11         816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
816       VAL       CG2       32.560       16.297       18.112       16.59         817       TRP       N       29.760       14.465       21.061       23.24         818       TRP       CA       28.891       14.345       22.223       25.45         819       TRP       C       29.223       13.143       23.061       25.99							
817       TRP N       29.760       14.465       21.061       23.24         818       TRP CA       28.891       14.345       22.223       25.45         819       TRP C       29.223       13.143       23.061       25.99							
818 TRP CA 28.891 14.345 22.223 25.45 819 TRP C 29.223 13.143 23.061 25.99							
819 TRP C 29.223 13.143 23.061 25.99					,	•	25.45
820 TRP O 29.403 13.249 24.269 25.20			C		13.143	-23 <del>.</del> 061	25.99
	820	TRP	0	29.403	13.249	24.269	25.20

821 822 823 824 825 826 827 828 829 830 831 832 833 834	TRP TRP TRP TRP TRP TRP TRP TRP GLU GLU GLU	CB CD1 CD2 NE1 CE2 CE3 CZ2 CZ3 CH2 N CA C	27.412 26.523 26.111 25.979 25.370 25.283 26.031 24.642 25.390 24.716 29.254 29.484 30.849 30.997	14.241 14.219 13.118 15.340 13.477 14.838 16.723 15.693 17.583 17.061 11.979 10.733 10.628 10.008	21.777 22.984 23.657 24.740 24.787 23.459 25.697 24.335 25.452 22.430 23.165 23.808 24.836	4 1 3 7 9 1 5 5 6 8	27.30 30.45 33.52 31.02 33.46 36.13 35.07 36.39 37.49 38.40 20.77 19.32 24.59 22.28
835 836	GLU GLU	CB CG	29.255 27.769	9.492 22.28 9.267 21.91		20.24 22.70	
837	GLU	CD	27.538	8.382 20.72		31.58	
838	GLU	OE1 OE2	28.422 26.263	7.828 20.07 8.280 20.450		28.77 22.97	
839 840	GLU GLN	N	31.853	11.228	23.20°		19.41
841	GLN	CA	33.216	11.131	23.689		18.18
842	GLN	C	33.531	12.204	24.662	2 .	20.10
843	GLN	0	34.595	12.205	25.267	7	21.48
844	GLN	CB	34.246	11.110	22.532	2	18.95
845	GLN	CG	33.923	9.996 21.53	2	19.84	•
846	GLN	CD	33.991	8.651 22.240		39.40	
847	GĿN	OE1	34.851	8.438 23.10		28.56	
848	GLN	NE2	33.079	7.739 21.94		26.05	40.00
849	LYS	N	32.598	13.105	24.794		19.33
	LYS	CA	32.677	14.184	25.768		18.87
851	LYS	C	33.789	15.161	25.514 26.422		22.27 19.41
852	LYS LYS	O CB	34.337 32.735	15.793 13.639	27.188		21.88
853 854	LYS	CG	31.398	13.126	27.65		19.83
855 .	LYS	CD	31.426	12.036	28.700		45.14
856	LYS	CE	30.008	11.554	29.070		76.91
857	LYS	NZ	29.545	10.345	28.337		96.48
858	SER	N	34.066	15.353	24.244	4	20.79
859	SER	CA	35.092	16.316	23.892	2	17.81
860	SER	С	34.624	17.729 .	24.20	5	21.92
861	SER	0	33.428	18.054	24.168		21.51
862	SER	CB	35.467	16.186	22.410		19.18
863	SER	OG	35.724	14.822	22.082		17.75
864	ARG	N	35.604	18.554	24.504		17.92
	ARG	-CA -		-19 <del>.</del> 949	24.80 23.62		16 <del>.</del> 44-
866	ARG	С	35.798	20.801	25.02	<i>f</i> .	13.30

867	ARG	0	35.360	21.920	23.437	19.40
868	ARG	CB	36.288	20.356 21.718	25.943 26.512	18.45 32.13
869 870	ARG ARG	CG CD	35.931 36.767	22.844	26.008	47.45
871	ARG	NE	36.397	24.110	26.645	78.03
872	ARG	CZ	35.147	24.537	26.840	72.70
873	ARG	NH1	34.080	23.841	26.491	57.33
874	ARG	NH2	34.975	25.713	27.412	58.72
875	GLY	N	36.733	20.313	22.843	18.99
876	GLY	CA	37.197	21.150	21.735	17.48
877	GLY	С	37.354	20.376	20.447	20.79
878	GLY	0	37.564	19.146	20.464	17.85
879	VAL	Ν	37.263	21.126	19.331	16.59
880	VAL	CA	37.432	20.590	17.992	13.96
881	VAL	С	38.479	21.445	17.339	19.19
882	VAL	.0	38.287	22.668	17.275	20.97
883	VAL	CB	36.129	20.735	17.192	15.20
884	VAL	CG1	36.421	20.354	15.742	14.08
885	VAL	CG2	35.021	19.821	17.752	15.12
886	VAL	N	39.576	20.856	16.870	13.97
887	VAL	CA	40.603	21.610	16.157	11.79
888	VAL	С	40.457	21.284	14.648	18.77
889	VAL	0	40.570	20.121	14.232	18.28
890	VAL	CB	41.958	21.180	16.675	12.96
891	VAL	CG1	43.076	21.854	15.888	14.53
892	VAL	CG2	42.053	21.591	18.138	13.41
893	MET	N ·	40.169	22.317	13.833	17.03
894	MET	CA	40.019	22.225	12.360	15.40
895	MET	C	41.205	22.846	11.677	18.09
896	MET	O	41.430	24.034	11.821	19.11 14.55
897	MET	CB	38.727 38.502	22.916 22.833	11.923 10.454	15.42
898	MET MET	CG SD	36.823	23.344	10.434	19.36
899 900	MET	CE	36.836	23.095	8.223 17.20	10.00
901	LEU	N	41.979	22.048	10.954	10.88
902	LEU	CA	43.193	22.584	10.379	10.33
903	LEU	C ·	43.143	22.854	8.877 13.92	10.00
904	LEU	Ö	44.145	23.245	8.271 14.65	
905	LEU	CB	44.326	21.595	10.679	13.36
906	LEU	CG '	44.519	21.273	12.178	16.29
907	LEU	CD1	45.594	20.155	12.274	14.38
908	LEU	CD2	45.006	22.542	12.913	13.63
909	ASN	N	41.968	22.712	8.311 13.55	
910	ASN	CA	41.826	22.925	6.905 17.04	
911-	-ASN-	C	40.701	23.897	6:601-18:91	·
912	ASN	O.	39.965	24.258	7.509 17.09	

913	ASN	СВ	41.343	21.584	6.310 18.53
914	ASN	CG	39.949	21.183	6.753 18.64
915	ASN	OD1	38.953	21.213	5.995 19.54
916	ASN	ND2	39.867	20.701	7.964 12.69
917	ARG	N	40.583	24.340	5.325 16.20
918	ARG	CA	39.429	25.199	4.934 18.31
919	ARG	C	38.419	24.316	4.226 23.61
920	ARG	Ö	38.770	23.219	3.769 20.22
921	ARG	CB	39.765	26.336	3.973 19.06
922	ARG	CG	40.782	27.259	4.610 27.14
923	ARG	CD	40.998	28.568	3.854 35.02
923	ARG	NE	41.400	28.489	2.457 76.50
925	ARG	CZ	42.181	27.557	1.889 100.00
	ARG	NH1	42.629	26.485	2.605 100.00
926		NH2	42.029	27.679	0.574 74.39
927	ARG			24.772	4.126 17.39
928	VAL	N C A	37.162	23.961	3.451 18.47
929	VAL	CÁ	36.162		1.972 24.72
930	VAL	С	36.529	23.752	1.424 20.57
931	VAL	0	36.433	22.657	
932	VAL	CB	34.781	24.569	3.651 20.83
933	VAL	CG1	33.815	24.070	2.560 18.89
934	VAL	CG2	34.319	24.222	5.084 18.54
935	MET	N	37.039	24.816	1.331 21.78
936	MET	CA	37.494	24.680	-0.05422.02
937	MET	С	39.008	24.831	-0.15423.31
938	MET	0	39.563	25.775	0.348 21.11
939	MET	CB	36.915	25.735	-0.97024.02
940	MET	CG	37.613	25.464	-2.29233.60
941	MET	SD	36.695	26.151	-3.672 42.56
942	MET	CE	35.122	25.238 °	-3.576 37.18
943	GLU	N	39.697	23.938	-0.824 18.74
944	GLU	CA	41.128	24.062	-0.945 18.84
945	GLU	С	41.474	23.440	-2.263 26.07
946	GLU	0	40.841	22.451	-2.688 29.43
947	GLU	CB	41.877	23.271	0.166 21.64
948	GLU	CG	41.562	23.754	1.595 24.84
949	GLU	CD	42.242	22.902	2.628 27.45
950	GLU	OE1	42.453	21.723	2.501 25.26
951	GLU	OE2	42.525	23.545	3.714 24.68
952	LYS	N	42.458	24.000	-2.90122.86
953	LYS	CA	42.899	23.487	-4.18723.50
954	LYS	С	41.742	23.348	-5.20429.15
955	LYS	0	41.767	22.481	-6.06130.13
956	LYS	СВ	43.790	22.226	-4.00926.67
957	LYS	- CG	45.143	22.628	-3.333-33.73
958	LYS	CD	46.022	21.518	-2.773 52.09
		-			

959 960	LYS LYS	CE NZ	47.424 48.324		22.009 22.031 24.167	,	-2.42239.14 -3.58176.66 -5.07623.79
961 962	GLY GLY	N CA	40.695 39.609		24.167		-6.01021.74
963	GLY	C	38.631		23.039		-5.69324.19
964	GLY	Ö	37.690		22.826		-6.457 29.23
965	SER	N	38.860		22.368		-4.620 16.90
966	SER	CA	37.939		21.336		-4.250 19.52
967	SER	С	37.336		21.506		-2.90122.34
968	SER	0	37.870		22.237		-2.07020.15
969	SER	CB	38.620		20.006		-4.177 25.21 5.539 47.60
970	SER	OG	38.845		19.720		-5.53847.69 -2.69315.65
971	LEU LEU	N CA	36.239 35.596		20.761 20.819		-1.374 14.31
972 973	LEU	C	36.183		19.731		-0.50221.78
974	LEU	O T	35.871		18.558		-0.677 26.91
975	LEU	СВ	34.104		20.610		-1.481 15.29
976	LEU	CG	33.470		21.756		-2.283 21.97
977	LEU	CD1	31.980		21.527		-2.467 21.98
978	LEU	CD2	33.677		23.043		-1.46826.21
979	LYS	Ν	37.037		20.122		0.436 17.45
980	LYS	CA	37.741		19.191		1.299 14.44
981	LYS	C	37.009		18.778		2.566 17.63 3.313 16.36
982	LYS	O	37.488 39.043		17.920 19.830		3.313 16.36 1.755 14.89
983 984	LYS LYS	ÇB CG	39.043		20.179		0.609 20.49
985	LYS	CD	40.109		19.031		-0.36521.67
986	LYS	CE	41.374		18.235		-0.21030.52
987	LYS	NZ	41.699		17.355		-1.37924.68
988	CYS	Ν	35.889		19.384		2.848 16.47
989	CYS	CA	35.155		19.078		4.070 16.96
990	CYS	С	33.741		19.677		3.961 19.21
991	CYS	0	33.563		20.642		3.244 18.82
992	CYS	CB	35.948		19.897		5.159 16.39
993	CYS	SG	35.331		19.703	•	6.887 18.71 4.700 16.15
994	ALA	N CA	32.760 31.415		19.154 19.714		4.739 14.64
995 996	ALA ALA	CA	31.413		20.957		5.690 19.23
997	ALA	Ö	32.233		21.145		6.585 17.35
998	ALA	СВ	30.467		18.651		5.231 13.19
999	GLN	N	30.461		21.869		5.475 21.44
1000	GLN	CA	30.365		23.045	,	6.336 24.00
1001	GLN	C	29.591		22.536		7.463 24.84
1002	GLN	0	28.375	•	22.686		7.394 24.39
1003	GLN	CB	29.530	_	24.166		5.663 26.72
1004	GLN	CG	29.391		25.440		6.544 22.81

1005	GLN	CD	30.676	25.983	7.104 25.37	
1003	GLN	OE1	30.793	26.198	8.315 30.89	
1007	GLN	NE2	31.616	26.296	6.240 21.26	
1007	TYR	NEZ	30.266	21.838	8.409 17.99	
		CA	29.535	21.156	9.487 15.44	
1009	TYR			21.150	10.748	22.11
1010	TYR	С	29.151		11.740	18.91
1011	TYR	0	28.643	21.358	9.883 16.63	10.91
1012	TYR	CB	30.286	19.866	10.522	17.61
1013	TYR	CG	31.599	20.198	9.749 16.52	17.01
1014	TYR	CD1	32.752	20.328		15 70
1015	TYR	CD2	31.683	20.328	11.910	15.79
1016	TYR	CE1	33.967	20.657	10.340	15.78
1017	TYR	CE2	32.904	20.620	12.511	12.70
1018	TYR	CZ	34.026	20.821	11.722	18.43
1019	TYR	ОН	35.226	21.088	12.310	18.28
1020	TRP	Ν	29.389	23.282	10.739	19.64
1021	TRP	CA	29.017	24.082	11.885	20.07
1022	TRP	C	28.335	25.344	11.398	20.86
1023	TRP -	0	28.609	25.785	10.296	18.14
1024	TRP	CB	30.276	24.405	12.737	19.79
1025	TRP	CG	31.146	25.427	12.111	20.56
1026	TRP	CD1	31.114	26.769	12.391	23.88
1027	TRP	CD2	32.177	25.255	11.089	19.89
1028	TRP	NE1	32.061	27.448	11.631	23.47
1029	TRP	CE2	32.720	26.554	10.813	23.19
1030	TRP	CE3	32.680	24.162	10.363	21.33
1031	TRP	CZ2	33.724	26.765	9.888 21.55	
1032	TRP	CZ3	33.681	24.396	9.418 21.71	
1033	TRP	CH2	34.190	25.681	9.187 22.67	
1034	PRO	N	27.472	25.933	12.244	19.83
1035	PRO	CA	26.755	27.155	11.892	19.59
1036	PRO	C	27.657	28.375	11.800	23.09
1037		·O .	28.534	28.630	12.627	24.72
1037	PRO	CB	25.736	27.390	13.013	20.41
	PRO	CG :	26.142	26.514	14.203	24.23
1039			27.223	25.561	13.671	19.47
1040	PRO	CD		29.188	10.787	20.72
1041	GLN	N	27.361	30.394	10.787	24.35
1042	GLN	CA	28.094		11.131	30.18
1043	GLN	C	27.360	31.604		
1044	GLN	0	27.958	32.680	11.285	28.65
1045	GLN	CB	28.440	30.572	9.138 25.68	
1046	GLN	CG	29.324	29.390	8.712 36.40	
1047	GLN	CD	29.769	29.566	7.304 56.84	-
1048	GLN	OE1	28.981	29.299	6.359 45.16	
1049	GLN	NE2		30.080	7.175 -51.88	04.00
1050	LYS	N	26.094	31.422	11.446	24.00

1051 1052 1053 1054 1055 1056 1057 1058 1059	LYS LYS LYS LYS LYS GLU	CA C O CB CG CD CE NZ N	25.374 24.547 23.907 24.653 23.416 23.256 24.577 25.063 24.612 23.924		32.529 32.118 31.078 33.411 32.791 33.389 33.985 33.363 32.988 32.760		12.050 13.199 13.158 11.123 10.569 9.195 1 8.716 1 7.463 1 14.210 15.437	00.0	Ò
1060 1061 1062 1063 1064	GLU GLU GLU GLU	C O CB	22.542 22.083 23.897 25.250		32.373 31.334 34.027 34.370		15.160 15.615 16.333 17.051		22.72 21.78 20.49 17.63
1065 1066 1067 1068	GLU GLU GLU	OE1 OE2 N	26.224 26.156 27.088 21.820		35.178 35.277 35.835 33.237		16.201 14.996 16.877 14.433		22.34 23.40 20.11 18.53
1069 1070 1071 1072 1073	GLU GLU GLU GLU	CA C O CB CG	20.418 20.107 18.937 19.614 20.050		32.935 31.763 31.405 34.172 34.671		14.214 13.281 13.088 13.817 12.449		19.99 27.09 23.53 20.56 21.72
1074 1075 1076 1077	GLU GLU GLU LYS	CD OE1 OE2 N	21.264 22.170 21.209 21.136	()	35.493 35.239 36.560 31.153	***	12.560 13.292 11.870 12.687		32.26 30.85 46.23 24.09
1078 1079 1080 1081 1082	LYS LYS LYS LYS	CA C O CB CG	20.821 21.462 22.539 21.378 20.579		30.032 28.780 28.413 30.287 31.355	•	11.815 12.346 11.875 10.448 9.738 5	3.04	25.40 32.06 34.08 25.92
1083 1084 1085 1086	LYS LYS LYS GLU	CD CE NZ N	20.028 18.577 18.161 20.846		30.875 30.432 29.607 28.148		8.415 6 8.487 8 7.339 1 13.337	1.11 1.13	24.74
1087 1088 1089 1090 1091	GLU GLU GLU GLU	CA C O CB CG	21.437 21.324 20.606 20.762 19.726		26.950 25.780 25.824 26.410 27.239	• .	13.921 12.983 12.017 15.183 15.866		24.02 28.33 25.06 25.15 51.90
1092 1093 1094	GLU GLU GLU MET	CD OE1 OE2	18.497 17.593 18.492 -21.986		27.376 26.572 28.538 -24.679	()	15.063 15.036 14.496 -13 <del>.</del> 350		40.35 36.72 33.99 24.61
1096	MET	CA	21.950		23.455		12.553		21.48

1097	MET	С	21.326	22.368	13.373	27.14
1098	MET	0	21.641	22.217	14.572	26.22
1090	MET	CB	23.369	22.980	12.072	21.43
	MET	CG	23.958	23.895	11.019	22.29
1100			25.666	23.460	10.592	26.20
1101	MET	SD				20.20
1102	MET	CE	25.256	22.169	9.435 23.94	27.52
1103	ILE	N	20.444	21.594	12.707	27.52
1104	ILE	CA	19.811	20.472	13.387	28.07
1105	ILE	С	20.179	19.186	12.668	32.13
1106	ILE	0	20.079	19.113	11.435	30.18
1107	ILE	CB	18.293	20.602	13.485	32.74
1108	ILE	CG1	17.977	21.495	14.686	32.67
1109	ILE	CG2	17.799	19.197	13.784	32.86
1110	ILE	CD1	16.777	22.374	14.453	40.14
1111	PHE	N	20.657	18.208	13.416	23.83
1112	PHE	CA	21.041	16.959	12.785	23.10
1113	PHE	C	19.998	15.956	13.154	25.20
1114	PHE	Ö	20.027	15.383	14.223	23.48
1115	PHE	СВ	22.477	16.491	13.147	22.44
1116		CG	23.457	17.603	12.869	21.09
	PHE			17.863	11.574	21.96
1117	PHE	CD1	23.901			
1118	PHE	CD2	23.914	18.415	13.901	20.96
1119	PHE	CE1	24.802	18.889	11.307	19.88
1120	PHE	CE2	24.819	19.449	13.666	21.74
1121	PHE	CZ	25.240	19.692	12.360	18.52
1122	GLU	N	19.041	15.794	12.276	26.50
1123	GLU	CA	17.949	14.903	12.589	27.86
1124	GLU	С	18.330	13.470	12.781	32.08
1125	GLU	0	17.727	12.809	13.608	35.43
1126	GLU	CB	16.877	14.981	11.517	30.62
1127	GLU	CG	16.580	16.453	11.155	62.94
1128	GLU	CD	15.389	16.595	10.252	100.00
1129	GLU	OE1	15.483	16.977	9.084 100.0	0
1130	GLU	OE2	14.265	16.211	10.846	100.00
1131	ASP	N ·	19.299	12.959	12.012	25.73
1132	ASP	CA	19.656	11.567	12.181	23.02
1133	ASP	C	20.185	11.281	13.545	28.07
		0	19.956	10.200	14.084	29.11
1134	ASP				11.107	25.54
1135	ASP	CB	20.632	11.086		
1136	ASP	CG	21.905	11.885	11.021	37.00
1137	ASP	OD1	22.084	12.993	11.515	30.66
1138	ASP	OD2	22.789	11.248	10.330	34.51
1139	THR	N	20.935	12.242	14.102	22.93
1140	THR	CA	21.496	12.004	15.402	20.67
	THR	C	20.850	12.755	16.525	23.97
1142	THR	0	21.319	12.650	17.645	24.14

1143	THR	СВ	23.011	12.160	15.438	24.59
1144	THR	OG1	23.323	13.466	15.015	22.89
1145	THR	CG2	23.629	11.120	14.521	20.72
1146	ASN	N	19.789	13.480	16.239	25.41
1147	ASN	CA	19.071	14.191	17.312	27.59
1148	ASN	С	19.850	15.245	18.085	29.45
1149	ASN	0	19.714	15.298	19.304	27.48
1150	ASN	CB	18.408	13.208	18.326	36.52
1151	ASN	CG	17.000	13.621	18.723	67.20
1152	ASN	OD1	16.346	14.422	18.030	53.22
1153	ASN	ND2	16.539	13.115	19.867	60.40
1154	LEU	Ν	20.633	16.084	17.377	24.61
1155	LEU	CA	21.440	17.164	17.988	23.07
1156	LEU	С	21.206	18.521	17.335	26.63
1157	LEU	0	21.001	18.634	16.126	23.32
1158	LEÜ	СВ	22.937	16.857	17.853	21.58
1159	LEU	CG	23.337	15.645	18.637	25.34
1160	LEU	CD1	24.640	15.072	18.051	25.68
1161	LEU	CD2	23.514	16.058	20.088	25.96
1162	LYS	N	21.305	19.557	18.160	23.08
1163	LYS	CA	21.182	20.874	17.655	21.49
1164	LYS	C	22.505	21.524	17.940	22.10
1165	LYS	Ō	23.066	21.305	18.982	23.53
1166	LYS	СВ	20.067	21.662	18.336	21.09
1167	LYS	CG	19.870	23.030	17.657	21.83
1168	LYS	CD	18.540	23.701	18.050	20.42
1169	LYS	CE	18.579	24.236	19.482	27.34
1170	LYS	NZ	17.233	24.626	19.969	28.39
1171	LEU	N	22.992	22.343	17.032	21.33
1172	LEU	CA	24.283	22.965	17.219	21.43
1173	LEU	C	24.163	24.452	16.940	23.20
1174	LEU	Ō	23.767	24.857	15.847	23.59
1175	LEU	СВ	25.209	22.322	16.142	22.80
1176	LEU	CG	26.646	22.855	16.136	23.00
1177	LEU	CD1	27.324	22.527	17.442	20.39
1178	LEU	CD2	27.437	22.271	14.965	24.15
1179	THR	N	24.498	25.276	17.874	20.33
1180	THR	CA	24.330	26.697	17.621	20.96
1181	THR	C ·	25.596	27.533	17.770	20.29
1182	THR	Ö	26.356	27.291	18.686	22.10
1183	THR	СВ	23.364	27.260	18.679	21.96
1184	THR	OG1	22.155	26.543	18.666	22.75
	THR	CG2	23.137	28.739	18.365	21.55
1186	LEU	N.	25.777	28.540	16.915	16.73
1187	LEU	CA.	26.914	29.399	17.070	-17.12
1188	LEU	C	26.594	30.412	18.199	25.18

1190 LI 1191 LI 1192 LI 1193 LI 1194 IL 1195 IL 1196 IL 1197 IL 1198 IL 1199 IL 1200 IL 1201 S 1204 S 1205 S 1204 S 1205 S 1206 G 1207 G 1210 G 1211 G 1212 G 1213 G 1214 G 1215 G 1216 G 1217 A 1218 A 1219 A 1221 A 1222 A 1223 A 1224 IL 1227 IL 1228 IL 1229 IL 1230 IL 1231 IL 1231 IL	E CA E E E E E E E E E E E E E E E E E E E		31.140 30.194 31.272 30.638 31.874 30.433 31.325 32.558 33.684 30.585 29.420 31.544 29.851 32.399 33.576 33.201 32.043 34.191 33.178 34.183 33.904 34.958 36.166 33.570 34.695 34.494 33.405 35.627 34.695 34.494 35.627 34.500 35.390 35.177 34.191 35.034 35.708 36.014 36.014 36.677 37.831 36.612 35.991 36.607	18.144 15.745 15.781 16.105 14.386 19.251 20.365 20.189 20.405 21.640 21.797 22.811 21.734 19.729 19.528 18.738 18.682 20.812 21.542 17.391 17.494 17.532 15.962 14.975 13.672 13.128 13.204 17.545 17.665 16.468 16.394 19.028 19.390 18.949 20.175 15.490 14.247 14.347 14.692 13.070 12.955 11.722 13.317	21.33 18.33 22.75 25.53 20.81 22.62 18.73 25.51 24.71 17.88 19.05 20.59 22.30 28.89 27.38 28.97 43.41 28.09 31.06 33.57 57.41 87.92 58.92 100.00 25.81 25.56 33.25 31.20 29.04 43.28 44.46 48.31 36.33 37.31 36.33 37.31
	E CD1 YS N YS CA	34.977 40.627 41.961	36.984 35.937 36.472	13.317 14.049 14.069	65.27 -22.86 - 23.68
1207 L		11.001	JJ. 1. 4		_5.55



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1235	LYS	С	42.448	36.470	12.610	25.39
1236	LYS	0	41.705	36.091	11.707	25.60
1237	LYS	CB	42.921	35.769	15.050	24.58
1238	LYS	CG	42.498	35.848	16.530	30.43
1239	LYS	CD	42.894	37.206	17.140	56.04
1240	LYS	CE	42.275	37.473	18.509	77.38
1241	LYS	NZ	43.040	38.439	19.321	100.00
1242	THR	N	43.669	36.907	12.395	22.88
1242	THR	CA	44.215	36.986	11.049	23.70
1243	THR	C	44.240	35.654	10.392	30.85
				35.565	9.210 34.77	30.00
1245	THR	0	43.898		10.942	20.00
1246	THR	CB	45.696	37.481	6 *	30.00
1247	THR	OG1	46.602	36.842	11.798	35.80
1248	THR	CG2	45.878	38.989	10.992	61.05
1249	TYR	N	44.743	34.649	11.108	21.60
1250	TYR	CA	44.882	33.376	10.456	18.56
1251	TYR	С	44.143	32.244	11.082	25.54
1252	TYR	0	44.324	31.078	10.663	28.06
1253	TYR	CB	46.334	33.048	10.304	18.65
1254	TYR	CG	46.986	32.802	11.597	20.73
1255	TYR	CD1	47.329	33.853	12.463	21.11
1256	TYR	CD2	47.327	31.498	11.937	20.37
1257	TYR	CE1	48.003	33.584	13.661	20.41
1258	TYR	CE2	47.992	31.216	13.130	21.74
		CZ	48.327	32.257	13.994	29.63
1259	TYR				15.159	23.15
1260	TYR	ОН	49.006	31.917		21.11
1261	TYR	N	43.337	32.562	12.101	
1262	TYR	CA	42.551	31.528	12.759	22.22
1263	TYR	С	41.350	32.139	13.375	24.48
1264	TYR	0	41.326	33.342	13.534	23.41
1265	TYR	CB	43.342	30.705	13.792	23.10
1266	TYR	CG	43.742	31.476	15.028	22.92
1267	TYR	CD1	44.930	32.194	15.073	20.83
1268	TYR	CD2	42.959	31.415	16.177	24.45
1269	TYR	CE1	45.287	32.851	16.247	21.19
1270	TYR	CE2	43.297	32.082	17.354	21.53
1271	TYR	CZ	44.483	32.785	17.378	21.56
1272	TYR	OH	44.815	33.471	18.512	22.47
1273	THR	N	40.369	31.311	13.688	20.51
1274	THR	CA,	39.136	31.768	14.317	20.65
1275	THR	C	38.791	30.762	15.378	25.40
		Ö	•	29.535	15.158	21.46
1276	THR		38.905		13.303	25.48
1277	THR	CB OC1	37.933	31.848		23.46
	THR-		38-230	32.801	12.320	
1279	THR	CG2	36.650	32.293	13.969	17.96
1280	VAL	N.	38.318	31.309	16.488	19.73

1281	VAL	CA	37.851	30.530	17.623	17.86
		C.	36.371	30.790	17.799	23.43
1282	VAL			31.937	17.908	23.75
1283	VAL	0	35.931			22.90
1284	VAL	CB	38.557	30.926	18.924	
1285	VAL	CG1	38.176	29.946	20.012	23.85
1286	VAL	CG2	40.057	30.808	18.737	23.85
1287	ARG	Ν	35.609	29.713	17.805	20.09
1288	ARG	CA	34.172	29.781	17.953	19.44
1289	ARG	С	33.699	29.002	19.174	23.12
1290	ARG	Ο.	34.205	27.901	19.557	20.13
1291	ARG	CB	33.431	29.182 <sup>\</sup>	16.745	16.93
1292	ARG	CG	33.792	29.938	15.473	24.10
1293	ARG	CD	33.027	29.483	14.215	26.79
1294	ARG	NE	33.620	30.035	12.964	35.76
1295	ARG	CZ	34.769	29.584	12.421	51.01
1296	ARG	NH1	35.509	28.558	12.930	52.65
1297	ARG	NH2	35.199	30.195	11.337	44.38
1298	GLN	N	32.711	29.625	19.799	16.92
1299	GLN	CA	32.040	28.948	20.858	20.12
		C		28.456	20.290	23.98
1300	GLN		30.679		19.711	23.75
1301	GLN	0	29.851	29.207		
1302	GLN	CB	31.889	29.796	22.106	23.97
1303	GLN	ĊG	31.076	29.015	23.134	41.20
1304	GLN	CD	30.559	29.958	24.176	67.03
1305	GLN	OE1	31.089	31.100	24.332	52.88
1306	GLN	NE2	29.502	29.485	24.841	56.56
1307	LEU	Ν	30.494	27.148	20.356	17.61
1308	LEU	CA	29.310	26.552	19.806	18.88
1309	LEU	С	28.563	25.874	20.920	26.36
1310	LEU	0	29.175	25.420	21.914	29.37
1311	LEU	СВ	29.714	25.412	18.853 · · ′	19.47
1312	LEU	CG	30.632	25.859	17.719 ~	24.78
1313	LEU	CD1	31.236	24.593	17.132	23.14
1314	LEU	CD2	29.825	26.634	16.632	19.35
1315	GLU	N	27.268	25.770	20.753	20.30
1316	GLU	CA ,	26.490	25.073	21.753	19.19
1317	GLU	C	25.861	23.866	21.128	23.29
1318	GLU	Ö	25.154	23.934	20.078	25.61
	GLU	CB	25.419	25.931	22.471	22.13
1319				25.130	23.502	24.76
1320	GLU	CG	24.553	26.017	23.909	45.63
1321	GLU	CD	23.408			51.61
1322	GLU	OE1	23.525	26.834	24.760	
1323	GLU	OE2	22.343	25.925	23.137	62.26
1324	LEU	N	26.125	22.751	21.814	21.06
1325	LEU	CA	25.635	21.497	21.378	22.31
1326	LEU	С	24.546	21.016	22.327	27.67

1354 1355 1356 1357 1358 1359 1360 1361 1362 1363 1364 1365 1366 1367 1368 1369	LUUUUUUUUUUUUUUU UN N N N N N N UUUUUUUU	O C C C D D C C O C C O C C O C C O C C C C	24.761 26.852 26.539 26.152 27.784 23.403 22.302 21.791 21.443 21.070 19.918 18.778 18.856 17.717 21.623 21.013 19.536 18.959 21.368 20.671 19.515 21.363 18.959 15.455 17.041 17.610 17.221 17.091 17.269 16.475 16.335 17.157 17.334 16.952 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631 17.522 17.631	20.942 20.528 19.163 19.220 18.312 20.711 20.214 18.886 18.671 21.580 22.616 21.235 18.021 16.727 17.002 17.366 15.757 14.311 13.396 16.255 16.484 17.273 18.466 18.478 19.717 15.169 14.567 14.567 14.663 14.986 16.882 17.266 13.086	23.522 21.268 20.645 19.149 20.815 21.735 22.466 21.998 20.844 22.382 23.287 23.311 22.768 23.997 22.728 22.721 23.883 23.812 23.355 21.535 21.535 22.278 22.392 20.108 17.959 20.120 22.738 23.285 24.335 24.355 25	26.01 24.03 23.92 20.97 23.65 25.02 26.18 33.77 34.24 28.09 31.05 37.23 35.49 35.58 36.35 44.05 29.96 30.39 44.87 46.33 28.60 34.74 39.55 49.55 50.59 71.99 84.31 74.28 74.29
1369 1370 1371	THR THR -GLN-	OG1 CG2 N	20.046 19.150 - 17.784	 15.266 13.086 17.191	 26.798 27.255 -26.081	63.24 59.51 -41.45
1372	GLN	CA	17.729	18.616	26.132	38.83

1373	GLN	С	18.894	19.1	29	26.948		39.20
1374	GLN	Ö	18.978	20.2		27.345		40.16
1375	GLN	СВ	16.416	19.1	91	26.628		41.08
1376	GLN	CG	15.319	19.2	28	25.576		61.17
1377	GLN	CD	14.099	19.9	68	26.091		89.86
1378	GLN	OE1	13.915	20.1	36	27.317		69.61
1379	GLN	NE2	13.273	20.4	37	25.155		100.00
1380	GLU	·N	19.813	18.2		27.212		35.40
1381	GLU	CA	21.012	18.6		27.881		35.65
1382	GLU	С	21.902	19.4		26.872		35.40
1383	GLU	0	21.846	19.3		25.623		30.83
1384	GLU	СВ	21.695	17.3		28.399		38.29
1385	GLU	CG	23.069	17.5		29.043		53.86
1386	GLU	CD	23.653	16.2		29.306		86.06
1387	GLU	OE1	22.927	15.1		29.234		47.15
1388	GLU	OE2	24.973	16.3		29.499		61.60
1389	THR	N	22.711	20.3		27.382 26.458		30.84 30.66
1390	THR	CA	23.466	21.1		26.816		36.12
1391	THR	С	24.932	21.1 21.1		28.021		33.20
1392	THR THR	O CB	25.281 22.790	21.1		26.411		39.57
1393 1394	THR	OG1	22.790	22.4		25.083		53.77
1394	THR	CG2	23.565	23.5		27.180		23.13
1396	ARG	N	25.775	21.2		25.751		30.39
1397	ARG	CA	27.226	21.3		25.923		29.24
1398	ARG	C	27.878	22.4		25.096		25.15
	ARG	Ö	27.469	22.6		23.981		24.36
1400	ARG	СВ	27.876	19.9		25.554		35.10
1401	ARG	CG	27.814	18.9		26.635		47.54
1402	ARG	CD	28.971	17.9		26.645		35.93
1403	ARG	NE	28.439	16.7	17	27.258		45.02
1404	ARG	CZ	28.503	15.5	48	26.702		70.35
1405	ARG	NH1	29.148	15.3	93	25.541		42.87
1406	ARG	NH2	27.9Ź2	14.5	22	27.339		59.02
1407	GLU	N	28.876	23.0	22	25.717		22.49
1408	GLU	CA	29.661	24.0		25.109		21.66
1409	GLU	С	30.904	23.4		24.481		24.49
1410	GLU	0	31.646	22.7		25.168		21.92
1411	GLU	CB	30.223	25.0		26.131		23.64
1412	GLU	CG	31.072	26.0		25.395		37.97
1413	GLU	CD	31.821	26.9		26.332		70.67
1414	GLU	OE1	31.279	27.5		27.207		100.00
1415	GLU	OE2	33.108	26.8		26.147	•	87.35
1416	ILE	N	31.145	23.7		23.201 22.541		20.06 19.28
1417	ILE	CA	32.319 33.071	23.1 24.3		21.942		22.12
1418	ILC		33.01 F	۷4.3	70	Z 1.34Z		££. 1£

	1419 1420	ILE	O CB	32.454 31.898	25.302 22.286	21.437 21.385	21.81 22.39
	1421	ILE	CG1	30.838	21.236	21.764	21.96
	1422	ILE	CG2	33.122	21.719	20.656	20.30
	1423	ILE	CD1	31.341	20.178	22.720	24.63.
	1424	LEU	N.	34.376	24.265	21.944	16.47
	1425	LEU	CA	35.137	25.326	21.321	16.94
	1426	LEU	C	35.661	24.796	20.041	21.55
	1427	LEU	Ö	36.126	23.648	19.980	20.21
	1428	LEU	СВ	36.341	25.704	22.198	18.75
	1429	LEU	CG	35.902	26.118	23.634	26.04
	1430	LEU	CD1	37.112	26.450	24.540	27.16
	1431	LEU	CD2	35.041	27.336	23.494	24.28
	1432	HIS	N	35.635	25.621	19.021	16.36
	1433	HIS	CA	36.118	25.183	17.703	14.50
	1434	HIS	C	37.275	26.073	17.333	20.44
	1435	HIS	Ö	37.118	27.293	17.360	20.81
	1436	HIS	СВ	34.956	25.388	16.717	15.22
	1437	HIS	CG	35.150	24.834	15.350	17.70
	1438	HIS	ND1	35.394	25.650	14.280	18.12
	1439	HIS	CD2	35.045	23.553	14.893	18.74
	1440	HIS	CE1	35.513	24.853	13.225	17.96
	1441	HIS	NE2	35.316	23.575	13.552	17.01
	1442	PHE	N	38.456	25.476	17.039	14.68
	1443	PHE	CA	39.642	26.223	16.705	14.36
	1444	PHE	C	39.905	25.964	15.252	18.76
	1445	PHE	.0	40.224	24.844	14.924	19.29
	1446	PHE	CB	40.854	25.765	17.540	14.14
	1447	PHE	CG	40.543	25.886	19.001	15.56
	1448	PHE	CD1	40.812	27.078	19.679	19.52
	1449	PHE	CD2	39.966	24.812	19.687	19.04
	1450	PHE	CE1	40.493	27.165	21.038	21.39
	1451	PHE	CE2	39.691	24.853	21.057	21.60
	1452	PHE	CZ	39.956	26.053	21.711	19.12
	1453	HIS	N	39.729	26.988	14.398	16.34
	1454	HIS	CA	39.850	26.842	12.954	16.68
	1455	HIS	С	41.036	27.540	12.393	22.59
	1456	HIS	0	41.056	28.776	12.342	20.03
	1457	HIS	CB	38.597	27.462	12.336	18.43
	1458	HIS	CG	38.504	27.092	10.899	22.05
	1459	HIS	ND1	37.487	27.555	10.113	21.32
	1460	HIS	CD2	39.322	26.311	10.139	23.08
	1461	HIS	CE1	37.665	27.073	8.892 20.85	
	1462	HIS	NE2	38.738	26.297	8.857 22.16	
	1463	TYR	-N	42.029	26.753	11.977	17.95
,	1464	TYR	CA	43.277	27.301	11.440	17.43
				*			

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1465	TYR	С	43.022	27.495	9.978 23.32	
1466	TYR	O.	42.787	26.542	9.283 19.33	
1467	TYR	СВ	44.414	26.250	11.602	16.22
1468	TYR	CG	45.848	26.801	11.601	19.15
1469	TYR	CD1	46.322	27.564	10.527	21.35
1470	TYR	CD2	46.732	26.525	12.639	19.74
1471	TYR	CE1	47.633	28.058	10.473	20.97
1472	TYR	CE2	48.053	26.987	12.600	20.96
1473	TYR	CZ	48.500	27.758	11.521	24.03
1474	TYR	OH	49.771	28.237	11.483	27.87
1475	THR	N	43.075	28.699	9.495 19.92	27.07
1475	THR	CA	42.722	28.877	8.099 22.16	
		C	43.882	29.154	7.134 28.45	
1477	THR		43.647	29.392	5.946 29.06	
1478	THR	0		30.007	8.008 25.85	
1479	THR	CB	41.656	31.210	8.473 23.68	
1480	THR	OG1	42.260	29.686	8.925 20.99	•
1481	THR	CG2	40.470		7.580 21.17	
1482	THR	N	45.126	29.138	6.619 20.89	
1483	THR	CA	46.174	29.416	6.556 28.24	
1484	THR	С	47.190	28.284	6.365 28.74	
1485	THR	0	48.385	28.506		
1486	THR	CB	46.906	30.686	7.023 27.18	
1487	THR	OG1	47.257	30.490	8.372 25.28	
1488	THR	CG2	46.033	31.944	6.834 22.60	
1489	TRP	N	46.743	27.029	6.778 21.40	
1490	TRP	CA	47.670	25.898	6.710 18.45	
1491	TRP	C	47.214	25.059	5.472 23.97	
1492	TRP	0	46.155	24.444	5.460 19.44	
1493	TRP	СВ	47.520	25.065	7.977 15.94	
1494	TRP	CG	48.522	23.957	8.065 15.20	
1495	TRP	CD1	49.281	23.420	7.059 17.12	•
1496	TRP	CD2	48.839	23.238	9.266 15.16	
1497	TRP	NE1	50.080	22.423	7.592 16.60	
1498	TRP	CE2	49.810	22.295	8.937 19.35	
1499	TRP	CE3	48.376	23.353	10.604	16.48
1500	TRP	CZ2	50.290	21.421	9.907 20.83	
1501	TRP	CZ3	48.843	22.520	11.569	17.13
1502	TRP	CH2	49.763	21.536	11.205	19.65
1503	PRO.	N	47.948	25.153	4.377 20.91	
1504	PRO	CA	47.535	24.503	3.150 19.93	
1505	PRO	C	47.609	22.996	3.191 21.72	
1506	PRO	0	48.534	22.409	3.802 19.55	
1507	PRO	CB	48.501	24.984	2.021 23.28	
1508	PRO	CG	49.570	25.796	2.717 28.22	
	-PRO-	CD	49.234	25.901	4.224 21.68	
1510	ASP	N	46.667	22.415	2.435 20.26	

1511 1512 1513 1514 1515 1516 1517 1518	ASP PHE	CA C O CB CG OD1 OD2 N	46.638 48.007 48.601 45.449 45.196 45.886 44.249 48.544	20.993 20.561 21.308 20.625 19.140 18.423 18.644 19.411	2.325 17.54 1.755 23.68 0.939 21.00 1.461 15.69 1.667 19.13 2.440 20.93 0.903 18.68 2.224 16.97
1519 1520 1521	PHE PHE PHE	CA C O	49.870 50.957 52.103	18.965 19.945 19.742	1.826 20.13 2.133 25.12 1.643 22.55
1522	PHE	CB	49.924	18.631	0.326 26.08
1523		CG	49.104	17.401	0.246 35.80
1524	PHE	CD1	49.297	16.474 17.211	1.282 48.33 -0.68140.42
1525	PHE	CD2 CE1	48.084 48.566	15.292	1.405 48.19
1526 1527		CE2	47.393	15.292	-0.63245.67
1528		CZ	47.618	15.075	0.407 46.93
1529		N	50.632	21.004	2.915 20.53
1530	GLY	CA	51.683	21.971	3.227 19.53
1531	GLY	C	52.004	22.104	4.708 21.83
1532	GLY	0	51.696	21.221	5.531 18.25
1533	VAL	N	52.661	23.217	5.033 18.02
1534	VAL	CA	53.016	23.487	6.390 16.83
1535	VAL	С	52.534	24.857	6.791 24.11
1536	VAL	0	52.177	25.662	5.955 24.27
1537	VAL	CB	54.512	23.437	6.545 20.78
1538	VAL	CG1	54.973	22.026	6.244 19.08
1539	VAL	CG2	55.079	24.462	5.559 22.82
1540	PRO	N	52.473 52.081	25.117 26.441	8.096 21.39 8.616 20.99
1541 1542	PRO PRO	CA C	53.080	27.518	8.157 26.47
1542	PRO	0 .	54.203	27.218	7.754 25.08
1544	PRO	CB	52.116	26.310	10.170 21.11
1545	PRO	CG	52.245	24.819	10.510 21.50
1546	PRO	CD	52.674	24.115	9.211 19.66
1547	GLU	N	52.696	28.777	8.213 24.24
1548	GLU	CA	53.593	29.831	7.749 23.61
1549	GLU	C	54.886	29.923	8.532 28.04
1550	GLU	Ο -	55.907	30.309	7.997 26.16
1551	GLU	СВ	52.881	31.191	7.690 25.09
1552	GLU	CG	51.548	31.090	6.895 64.56
1553	GLU	CD OF1	50.479	32.181	7.111 100.00 8.113 79.43
1554	GLU	OE1	49.716	32.227	6.070 91.77
1555 1556	GLU SER	OE2	50.381 54.859	33.007 29.633	9.821 22.79
1330	SER	IN:	JT.0J8	20.000	U.UL 1 LL.1U

1557	SER	CA	56.080	29.729	10.605	20.38
1558	SER	C ·	55.893	28.889	11.822	23.57
1559	SER	Ö	54.788	28.588	12.204	24.21
1560	SER	СВ	56.352	31.159	11.079	23.98
1561	SER	OG	55.221	31.636	11.819	21.35
1562	PRO	N	56.970	28.495	12.436	22.91
1563	PRO	CA .	56.805	27.775	13.649	23.08
1564	PRO	C	56.050	28.657	14.655	26.93
1565	PRO	Ö	55.238	28.194	15.396	25.45
1566	PRO	СВ	58.230	27.505	14.170	24.65
1567	PRO	CG	59.143	27.550	12.965	28.02
1568	PRO	CD	58.397	28.442	11.979	23.89
1569	ALA	N	56.300	29.973	14.661	23.54
1570	ALA	CA	55.629	30.881	15.613	21.90
1571	ALA	C	54.118	30.891	15.479	20.92
1572	ALA	Ö	53.348	30.851	16.457	19.69
1573	ALA	СВ	56.248	32.309	15.569	21.00
1574	SER	N	53.675	30.917	14.234	20.18
1575	SER	CA	52.201	30.957	14.031	21.57
1576	SER	C	51.539	29.677	14.476	24.10
1577	SER	Ö	50.457	29.670	15.047	23.15
1578	SER	СВ	51.769	31.348	12.606	25.87
1579	SER	OG	52.780	31.001	11.688	40.00
1580	PHE	N	52.229	28.573	14.201	19.09
1581	PHE	CA	51.685	27.283	14.553	15.85
1582	PHE	C	51.650	27.200	16.075	21.97
1583	PHE	Ö	50.659	26.814	16.684	20.90
1584	PHE	CB	52.611	26.166	13.984	18.25
1585	PHE	CG	52.293	24.814	14.607	18.09
1586	PHE	CD1	51.234	24.063	14.104	16.30
1587	PHE	CD2	53.032	24.327	15.687	19.26
1588	PHE	CE1	50.907	22.860	14.722	18.09
1589	PHE	CE2	52.685	23.149	16.344	23.27
1590	PHE	CZ	51.600	22.423	15.850	21.69
1591	LEU	N	52.774	27.544	16.720	20.33
1592	LEU	CA	52.866	27.418	18.177	19.90
1593	LEU	C	51.888	28.318	18.868	23.96
1594	LEU	Ö	51.233	27.949	19.866	23.17
1595	LEU	СВ	54.290	27.689	18.661	20.12
	LEU	CG	55.212	26.537	18.324	22.21
1596 1597	LEU	CD1	56.679	26.965	18.499	22.08
	LEU	CD1	54.843	25.331	19.215	23.19
1598		N		29.520	18.303	22.18
1599	ASN ASN	CA	51.778 50.801	30.490	18.835	21.35
1600		CA	49.403	29.847	18.865	21.98
1601	ASN		48.693	29.047	19.870	21.88
1602	ASN	0	40.033	25.320	19.070	21.00

1603 1604	ASN ASN	CB CG	50.785 49.807	31.816 32.839	18.027 18.619 19.783	20.82 24.28
1605	ASN	OD1 ND2	49.974 48.763	33.256 33.220	17.844	21.90 19.52
1606 1607	ASN PHE	ND2 N	49.044	29.156	17.743	18.21
1608	PHE	CA	47.745	28.501	17.628	14.93
1609	PHE	C	47.666	27.300	18.571	19.31
1610.	PHE	0	46.638	27.092	19.233	19.07
1611	PHE	CB	47.530	28.099	16.167	15.26
1612	PHE	CG	46.276	27.312	15.946	19.10
1613	PHE	CD1	45.072	27.942	15.602	22.25
1614	PHE	CD2	46.280	25.910	16.038	14.28
1615	PHE	CE1	43.887	27.210	15.433	20.75
1616	PHE	CE2	45.122	25.167	15.823	12.24
1617	PHE	CZ	43.919	25.815	15.533	13.08
1618	LEU	N	48.750	26.511	18.662	18.17
1619	LEU	CA	48.729	25.368	19.546	17.98
1620	LEU	C	48.499	25.780	21.000	21.46
1621	LEU	0	47.707	25.217	21.756	18.26
1622	LEU	СВ	50.052	24.635	19.403	18.33
1623	LEU	CG	50.151	23.515	20.450	21.15
1624	LEU	CD1	51.503	22.791	20.311	23.59
1625	LEU	CD2	48.978	22.520	20.369	15.46
1626	PHE	N	49.225	26.808	21.416	20.08
1627	PHE	CA	49.049	27.353	22.789	21.32
1628	PHE	С	47.628	27.968	23.006	24.42
1629	PHE	Ο .	47.052	27.870	24.091	24.60
1630	PHE	CB	50.175	28.286	23.244	20.93
1631	PHE	CG	51.389	27.451	23.568	25.75
1632	PHE	CD1	51.876	26.502	22.666	24.29
1633	PHE	CD2	52.059	27.590	24.788	33.03
1634	PHE	CE1	52.958	25.669	22.967	25.19
1635	PHE	CE2	53.152	26.773	25.100	36.27
1636	PHE	CZ	53.595	25.799	24.198	30.54
1637	LYS	N	47.012	28.564	21.981	19.37
1638	LYS	CA	45.650	29.029	22.169	19.01
1639	LYS	C	44.755	27.830	22.565	23.16
1640	LYS	0	43.965	27.875	23.510	20.57
1641	LYS	CB	45.064	29.654	20.900	20.68
1642	LYS	CG	44.966	31.167	21.019	54.23
1643	LYS	CD	43.679	31.660	21.671	58.81
1644	LYS	CE	43.329	33.110	21.340	79.60
1645	LYS	NZ	42.204		20.403	90.92
1646	VAL	N	44.903	26.725	21.833	18.90
1647	VAL	CA	44.143	25.527	22.117	17.53
1648	VAL	С	44.447	25.027	23.508	21.15

1649	VAL	0	43.558	24.712	24.284	19.52
1650	VAL	СВ	44.432	24.450	21.047	17.34
1651	VAL	CG1	43.692	23.154	21.348	16.63
1652	VAL	CG2	44.052	24.997	19.628	15.16
1653	ARG	N	45.712	24.939	23.832	20.55
1654	ARG	CA	46.047	24.444	25.161	19.47
1655	ARG	C	45.455	25.327	26.259	21.39
1656	ARG	Ö	44.954	24.911	27.288	21.16
1657	ARG	СВ	47.560	24.512	25.353	17.77
	ARG	CG	48.312	23.372	24.672	24.26
1658				23.572	24.620	22.42
1659	ARG	CD	49.824		24.419	26.10
1660	ARG	NE	50.439	22.292	25.113	40.08
1661	ARG	CZ	51.464	21.787		
1662	ARG	NH1	52.063	22.472	26.095	23.77
1663	ARG	NH2	51.909	20.558	24.796	21.91
1664	GLU	N ,	45.604	26.581	26.047	20.31
1665	GLU	CA	45.141	27.497	27.028	21.25
1666	GLU	С	43.668	27.493	27.214	26.86
1667	GLU	0	43.228	27.801	28.314	30.19
1668	GLU	CB	45.646	28.874	26.753	23.38
1669	GLU	CG	47.087	28.893	27.173	37.92
1670	GLU	CD	47.736	30.184	26.883	65.18
1671	GLU	OE1	47.282	30.979	26.063	63.04
1672	GLU	OE2	48.853	30.321	27.578	59.99
1673	SER	N	42.897	27.123	26.213	20.78
1674	SER	CA	41.438	27.087	26.408	20.46
1675	SER	Ċ	41.009	26.041	27.387	30.52
1676	SER	0	39.864	25.951	27.744	37.57
1677	SER	СВ	40.714	26.707	25.130	17.75
1678	SER	OG	40.998	25.358	24.799	21.13
1679	GLY	N	41.850	25.128	27.755	27.53
1680	GLY	CA	41.324	24.128	28.636	24.97
1681	GLY	C	40.817	22.896	27.894	35.66
1682	GLY	Ö	40.571	21.813	28.504	38.98
1683	SER	N	40.733	23.002	26.556	26.31
1684	SER	CA	40.241	21.873	25.787	23.11
1685	SER	C.	41.002	20.591	25.925	33.29
.1686	SER	0	40.402	19.563	25.614	33.64
1687		CB	40.402	22.177	24.303	24.92
	SER		39.263	23.274	24.065	24.93
1688	SER	OG			26.294	30.38
1689	LEU	N	42.318	20.630		30.40
1690	LEU	CA	43.103	19.378	26.359	38.35
1691	LEU	C	43.108	18.701	27.742	
	LEU	0	43.678	17.631	27.992	38.03
1693	LEU	CB	44.513	19.515	25.757	29.26
1694	LEU	CG	44.530	20.316	24.461	31.07

1695	LEU	CD1	45.948	20.591	24.014	27.31
1696	LEU	CD2	43.820	19.572	23.354	34.10
1697	SER	N	42.463	19.367	28.644	34.73
1698	SER	CA	42.363	18.870	29.968	34.22
1699	SER	C	41.847	17.426	30.022	36.01
1700	SER	Ŏ	40.925	17.051	29.369	31.19
1701	SER	СB	41.501	19.801	30.780	39.43
1702	SER	ÖĞ	41.658	19.437	32.131	55.74
1703	PRO	N	42.456	16.621	30.848	39.48
1704	PRO	CA	42.093	15.228	31.048	39.86
1705	PRO	C	40.730	15.105	31.735	38.86
1706	PRO	Ö	40.123	14.050	31.847	37.15
1707	PRO	СВ	43.162	14.696	31.998	42.88
1708	PRO	CG	43.756	15.909	32.728	48.04
1709	PRO	CD	43.253	17.145	31.996	43.33
1710	GLU	N ·	40.212	16.194	32.214	32.75
1711	GLU	CA	38.901	16.068	32.791	33.48
1712	GLU	C	37.828	16.086	31.693	31.63
1713	GLU	Ö	36.656	16.014	31.931	31.89
1714	GLU	CB	38.647	17.159	33.840	37.32
1715	GLU	CG	38.309	18.559	33.240	65.45
1716	GLU	CD	39.318	19.626	33.589	92.79
1717	GLU	OE1	40.366	19.358	34.151	100.00
1718	GLU	OE2	39.027	20.832	33.145	79.45
1719	HIS	N	38.217	16.238	30.459	23.47
1720	HIS	CA	37.246	16.264	29.407	21.41
1721	HIS	C	37.579	15.102	28.544	24.76
1722	HIS	Ö	38.664	14.584	28.727	24.90
1723	HIS	СВ	37.581	17.439	28.476	23.84
1724	HIS	CG	37.356	18.755	29.102	29.53
1725	HIS	ND1	36.121	19.085	29.633	32.52
1726	HIS	CD2	38.191	19.824	29.252	34.49
1727	HIS	CE1	36.203	20.348	30.081	33.65
1728	HIS	NE2	37.441	20.813	29.875	35.22
1729	GLY	N	36.738	14.774	27.539	20.61
1730	GLY	CA	37.086	13.704	26.621	18.66
1731	GLY	С	38.225	14.238	25.771	24.63
1732	GLY	0	38.659	15.427	25.902	26.39
1733	PRO	N	38.720	13.391	24.886	20.23
1734	PRO	CA	39.824	13.749	23.973	16.08
1735	PRO	С	39.406	14.853	23.015	16.63
1736	PRO	O	38.315	14.885	22.519	21.70
1737	PRO	СВ	40.065	12.516	23.099	18.85
1738	PRO	CG	39.214	11.419	23.667	23.51
1739		-CD	38.245	12.018	24.690	21.40
1740	VAL	N	40.277	15.758	22.729	17.27
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	1741	VAL	CA	39.952	16.799	21.788	16.61
	1742	VAL	С	39.821	16.100	20.459	20.32
	1743	VAL	0	40.473	15.057	20.168	20.45
•	1744	VAL	CB	41.125	17.820	21.737	20.35
	1745	VAL	CG1	42.315	17.135	21.117	22.42
•	1746	VAL	CG2	40.889	19.002	20.793	20.42
•	1747	VAL	N	38.972	16.637	19.619	16.18
•	1748	VAL	CA	38.847	16.060	18.263	13.46
•	1749	VAL	С	39.700	16.901	17.328	20.85
•	1750	VAL	0	39.635	18.187	17.316	22.85
•	1751	VAL	CB	37.396	16.152	17.783	16.77
•	1752	VAL	CG1	37.337	15.760	16.317	16.82
•	1753	VAL	CG2	36.566	15.164	18.549	16.12
•	1754	VAL	N	40.549	16.250 <sub>.</sub>	16.565	16.13
•	1755	VAL	CA	41.389	17.029	15.642	14.73
	1756	VAL	С	41.154	16.603	14.236	19.68
	1757	VAL	0	41.111	15.408	13.968	18.53
	1758	VAL	CB	42.844	16.737	15.897	16.28
•	1759	VAL	CG1	43.703	17.511	14.901	17.56
	1760	VAL	CG2	43.184	17.097	17.349	17.81
•	1761	HIS	N	40.996	17.532	13.289	15.11
	1762	HIS	CA	40.798	17.040	11.911	12.07
	1763	HIS	С	41.283	18.025	10.893	18.30
	1764	HIS	0	41.478	19.204	11.185	18.91
٠.	1765	HIS	CB	39.337	16.682	11.545	13.93
٠	1766	HIS	CG	38.478	17.919	11.267	15.77
	1767	HIS	ND1	38.367	18.469	9.979 15.86	
	1768	HIS	CD2	37.681	18.652	12.088	16.69
•	1769	HIS	CE1.	37.560	19.518	10.045	16.85
	1770	HIS	NE2	37.137	19.670	11.291	18.34
	1771	CYS	Ν	41.470	17.478	9.681 16.12	
	1772	CYS	CA	41.899	18.201	8.547 13.36	
	1773	CYS	С	40.993	17.660	7.461 17.96	
	1774	CYS	0	39.874	17.322	7.714 14.94	
	1775	CYS	CB	43.356	18.020	8.128 -12.46	4.
	1776	CYS	SG	44.095	16.346	8.360 18.95	
	1777	SER	N	41.479	17.548	6.218 15.33	
	1778	SER	CA	40.606	16.949	5.208 15.04	
	1779	SER	С	40.523	15.407	5.439 18.57	
	1780	SER	0	39.439	14.799	5.469 17.07	
	1781	SER	CB	41.042	17.237	3.766 14.15	
	1782	SER	OG	40.022	16.672	2.921 17.05	
	1783	ALA	Ν.	41.695	14.781	5.624 14.98	
	1784	ALA	CA	41.676	13.317	5.798 13.26	
,	1785	ALA	.C	41.900	12.903	7.242 18.49	
	1786	ALA	0	41.702	11.727	7.613 17.47	

1787 1788 1789 1790 1791 1792 1793 1794 1795 1796	ALA GLY GLY GLY ILE ILE ILE ILE	CB N CA C O N CA C O CB	42.793 42.343 42.600 43.930 44.071 44.929 46.226 47.402 48.386 46.405	12.703 13.867 13.538 12.809 11.993 13.082 12.446 13.347 13.343 11.141	4.988 14.41 8.075 16.70 9.471 15.45 9.660 18.39 10.600 8.789 13.74 8.919 11.49 8.720 18.18 9.475 19.37 8.141 16.27	16.53
1797 1798	ILE	CG1 CG2	46.367 45.308	11.366 10.153	6.605 17.55 8.566 15.49	
1799	IĽE	CD1	46.493	10.057	5.781 14.58	
1800	GLY	Ν	47.379	14.148	7.683 16.11	
1801	GLY	CA ,	48.606	14.939	7.449 16.70	
1802	GLY	С	48.766	16.139	8.371 19.48	
1803	GLY	0	49.669	16.167	9.200 20.37	
1804	ARG	N	47.898	17.159	8.203 14.27	1
1805	ARG	CA	47.981	18.302	9.086 14.68	17 72
1806	ARG	C	47.634	17.835	10.502	17.73
1807	ARG	0	48.295	18.218	11.470 8.599 12.68	19.43
	ARG	CB	47.114	19.444 20.033	8.599 12.68 7.287 13.57	
1809	ARG	CG CD	47.672 46.645	20.033	6.671 11.34	
1810 1811	ARG ARG	NE	45.594	20.360	5.930 16.97	
1812	ARG	CZ	44.676	20.854	5.129 31.69	
1813	ARG	NH1	44.618	22.185	4.971 16.84	
1814	ARG	NH2	43.766	20.109	4.490 15.94	•
1815	SER	N	46.626	16.986	10.647	15.68
1816	SER	CA	46.298	16.517	12.015	14.86
1817	SER	C	47.460	15.831	12.671	17.12
1818	SER	Ō	47.726	16.037	13.863	16.53
1819	SER	СВ	45.169	15.510	11.989	17.35
1820	SER	OG	44.028	16.136	11.424	17.53
1821	GLY	N	48.157	14.979	11.913	15.45
1822	GLY	CA	49.301	14.268	12.486	11.73
1823	GLY	С	50.394	15.213	12.935	17.63
1824	GLY	0	51.056	15.041	13.964	17.72
1825	THR	Ν	50.627	16.231	12.100	16.30
1826	THR	CA	51.651	17.225	12.377	15.64
1827	THR	С	51.360	17.997	13.664	19.70
1828	THR	0	52.171	18.184	14.543	17.95
1829	THR	CB	51.794	18.168	11.171	23.66
1830	THR	OG1	52.110	17.428	9.987 19.58	10.00
	THR.	CG2	52.870	19.217	11.433	19.33
1832	PHE	N	50.158	18.420	13.807	15.63

1833	PHE	CA	49.780	19.140	14.983	15.63
1834	PHE	C	49.909	18.259	16.248	20.88
1835	PHE	Ö	50.425	18.670	17.305	20.45
1836	PHE	СВ	48.260	19.563	14.772	17.22
1837	PHE	CG	47.593	20.201	15.997	17.55
1838	PHE	CD1	47.575	21.585	16.155	17.33
1839	PHE	CD2	46.967	19.435	16.983	17.40
1840	PHE	CE1	46.983	22.172	17.278	17.10
1841	PHE	CE2	46.361	20.001	18.114	18.42
1842	PHE	CZ	46.365	21.392	18.257	14.40
		N	49.390	17.033	16.193	17.18
1843	CYS			16.201	17.366	14.77
1844	CYS	CA	49.432		17.694	21.33
1845	CYS	C	50.843	15.806	18.842	20.61
1846	CYS	0	51.225	15.694	17.191	18.22
1847	CYS	CB	48.573	14.933		
1848	CYS	SG	46.804	15.307	17.081	23.37
1849	LEU	N	51.643	15.520	16.693	19.44
1850	LEU	CA	52.998	15.101	17.015	18.15
1851	LEU	С	53.714	16.200	17.797	19.49
1852	LEU	0	54.425	15.954	18.786	19.58
1853	LEU	CB	53.792	14.823	15.705	16.40
1854	LEU	CG.	55.288	14.537	15.965	18.63
1855	LEU	CD1	55.482	13.249	16.751	16.35
1856	LEU	CD2	55.942	14.240	14.617	19.47
1857	ALA	Ν	53.571	17.449	17.313 ·	16.69
1858	ALA	CA	54.249	18.534	18.012	. 15.40
1859	ALA .	С	53.697	18.630	19.431	19.18
1860	ALA	0	54.419	18.775	20.406	16.89
1861	ALA	CB	54.115	19.859	17.273	14.90
1862	ASP	N	52.390	18.527	19.557	19.72
1863	ASP	CA	51.781	18.658	20.893	17.54
1864	ASP	С	52.338	17.650	21.887	20.70
1865	ASP	0	52.697	17.949	23.038	19.79
1866	ASP	СВ	50.237	18.552	20.829	15.42
1867	ASP	CG	49.674	18.953	22.159	24.54
1868	ASP	OD1	50.072	19.914	22.820	19.87
1869	ASP	OD2	48.833	18.076	22.624	22.29
1870	THR	N	52.385	16.427	21.396	16.78
1871	THR	CA	52.782	15.309	22.210	14.97
1872	THR	C	54.217	15.416	22.614	18.91
1873	THR	Ö	54.563	15.159	23.809	16.57
1874	THR	СВ	52.438	13.929	21.591	19.52
1875	THR	0G1	51.035	13.752	21.566	22.01
1876	THR	CG2	53.027	12.780	22.446	15.17
1877	CYS	N	-55-074	15.710 -	21.605	16.93
1878	CYS	CA	56.498	15.818	21.947	16.87
1070		<b>5</b> / (	30.400	10.010		

1879	CYS	С	56.751	16.927	22.968	20.74
1880	CYS	0	57.647	16.825	23.812	19.23
1881	CYS	CB	57.335	16.123	20.706	18.21
1882	CYS	SG	57.376	14.721	19.578	21.43
1883	LEU	N	55.983	17.999	22.886	17.24
1884	LEU	CA	56.208	19.110	23.805	16.36
1885	LEU	С	55.758	18.707	25.185	22.13
1886	LEU	0	56.343	19.082	26.201	21.35
1887	LEŲ	CB	55.501	20.411	23.335	15.49
1888	LEU	CG	56.258	21.020	22.155	16.90
1889	LEU	CD1	55.474	22.152	21.560	18.67
1890	LEU	CD2	57.642	21.489	22.581	19.42
1891	LEU	Ν	54.709	17.914	25.244	18.58
1892	LEU	CA	54.187	17.454	26.563	19.33
1893	LEU	С	55.127	16.424	27.211	20.57
1894	LEU	0	55.403	16.398	28.406	17.16
1895	LEU	CB	52.758	16.888	26.411	20.29
1896	LEU	CG	52.083	16.494	27.720	23.78
1897	LEU	CD1	51.799	17.757	28.557	22.08
1898	LEU	CD2	50.756	15.824	27.386	26.13
1899	LEU	N	55.661	15.528	26.417	20.05
1900	LEU	CA	56.607	14.567	27.014	22.38
1901	LEU	С	57.798	15.295	27.539	21.44
1902	LEU	0	58.333	14.922	28.548	18:07
1903	LEU	CB	57.204	13.588	25.971	23.67
1904	LEÜ	CG	56.223	12.489	25.668	30.34
1905	LEU	CD1	56.535	11.952	24.272	35.67
1906	LEU	CD2	56.358	11.418	26.732	23.09
1907	MET	N <sup>*</sup>	58.247	16.315	26.802	17.46
1908	MET	CA	59.411	17.026	27.246	21.88
1909	MET	C .	59.093	17.791	28.540	23.63
1910	MET	0	59.925	17.939	29.443	21.17
1911	MET	СВ	60.016	17.837	26.046	26.35
1912	MET	CG	60.123	19.321	26.201	31.09
1913	MET	SD	61.319	20.100	25.086	31.12
1914	MET	CE	61.218	18.941	23.667	24.10
1915	ASP	N	57.837	18.250	28.614°	21.37
1916	ASP	CA	57.357	18.992	29.777	19.58
1917	ASP	С	57.355	18.111	31.012	26.30
1918	ASP	0	57.669	18.490	32.147	25.46
1919	ASP	СВ	55.936	19.493	29.486	18.02
1920	ASP	CG	55.662	20.778	30.190	23.00
1921	ASP	OD1	56.518	21.337	30.864	26.14
1922	ASP	OD2	54.428	21.203	30.067	21.46
1923		N	56.941	16.910	30.809	26.75
1924	LYS	CA	56.835	15.971	31.911	27.98
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1925	LYS	С	58.142	15.701	32.588	30.57
1926	LYS	Ö	58.233	15.653	33.824	29.41
1927	LYS	CB	56.113	14.673	31.511	31.11
1928	LYS	CG	56.050	13.703	32.679	62.81
1929	LYS	CD	54.796	12.831	32.771	87.49
1930	LYS	CE	54.830	11.905	34.006	100.00
1931	LYS	NZ	54.107	10.613	33.862	100.00
1932	ARG	N	59.175	15.573	31.790	29.07
		CA	60.454	15.266	32.352	31.74
	ARG		61.458	16.356	32.286	38.39
1934	ARG	С		16.350	32.699	42.73
1935	ARG	0	62.576		31.574	42.79
1936	ARG	CB	61.007	14.104		44.98
1937	ARG	CG	60.626	14.244	30.128	
1938	ARG	CD	61.065	13.064	29.267	65.93
1939	ARG	NE	60.195	11.933	29.445	74.57
1940	ARG	CZ	59.885	10.983	28.560	88.88
1941	ARG	NH1	60.356	10.889	27.280	42.03
1942	ARG	NH2	59.037	10.075	29.022	91.14
1943	LYS	Ν	61.090	17.490	31.753	28.22
1944	LYS	CA	62.065	18.538	31.662	25.40
1945	LYS	С	63.354	18.059	31.041	28.13
1946	LYS	0	64.435	18.477	31.438	25.94
1947	LYS	CB	62.253	19.262	32.958	26.58
1948	LYS	CG	60.936	19.905	33.457	27.81
1949	LYS	CD	60.409	21.051	32,575	14.64
1950	LYS	CE	59.256	21.768	33.219	19.71
1951	LYS	NZ	58.583	22.690	32.312	24.14
1952	ASP	N	63.240	17.207	30.024	22.76
1953	ASP	CA	64.428	16.721	29.406	22.09
1954	ASP	С	64.228	16.564	27.898	30.50
1955	ASP	Ō	63.820	15.533	27.405	33.52
1956	ASP	СВ	64.769	15.377	30.068	24.25
1957	ASP	CG	65.984	14.741	29.416	34.47
1958	ASP	OD1	66.675	15.328	28.608	33.87
1959	ASP	OD2	66.182	13.489	29.778	42.04
1960	PRO	N	64.523	17.584	27.151	24.81
1961	PRO	CA	64.355	17.595	25.725	25.70
1962	PRO	·C	65.131	16.551	24.997	31.12
1963	PRO	Ö	64.707	16.018	23.971	31.92
1964	PRO	СВ	64.832	18.944	25.251	29.74
1965	PRO	CG	64:947	19.803	26.511	32.73
1966	PRO	CD	65.066	18.845	27.678	26.52
1967	SER	N-	66.264	16.244	25.538	29.07
1967	SER	CA	67.077	15.273	24.890	31.78
	SER	CA	66.479	13.911	24.933	37.92
1969				13.023	24.955	45.55
1970	SER	0	66.793	13.023	24. IJU	+∪.∪∪

1971	SER	СВ	68.530	15.309	25.38		42.27
1972	SER	OG	69.183	16.449	24.808		59.55
1973	SER	N	65.580	13.735	25.830		28.18
1974	SER	CA	64.970	12.446	25.947		28.28
1975	SER	С	63.888	12.202	24.886		32.74
1976	SER	0	63.293	11.107	24.796		34.35
1977	SER	CB	64.304	12.317	27.328	3	30.87
1978	SER ·	OG	63.068	13.050	27.35	7	38.21
1979	VAL	N	63.544	13.269	24.19°	1	26.20
1980	VAL	CA	62.453	13.223	23.204	4	23.76
1981	VAL	С	62.902	12.743	21.809	9	26.52
1982	VAL	0	63.645	13.435	21.074	4	28.45
1983	VAL	CB	61.667	14.568	23.186	3	25.58
1984	VAL	CG1	60.555	14.594	22.126	3	25.16
1985	VAL	CG2	61.053	14.793	24.553	3	24.66
1986	ASP	Ň	62.404	11.567	21.409	9	21.57
1987	ASP	CA	62.732	11.009	20.11	1	19.61
1988	ASP	C	61.521	11.154	19.182	2	26.16
1989	ASP	Ö	60.569	10.318	19.178	3	26.94
1990	ASP	СВ	63.073	9.533 20.32	0	22.22	
1991	ASP	CG	63.559	8.879 19.07		34.42	
1992	ASP	OD1	63.392	9.325 17.93		28.99	
1993	ASP	OD2	64.139	7.760 19.35		40.03	
1994	ILE	N	61.550	12.245	18.38	1	23.48
1995	ILE	CA	60.414	12.608	17.503		19.92
1996	ILE	C	59.899	11.503	16.616		24.71
1997	ILE	Ö	58.701	11.191	16.558		22.68
1998	ILE	CB	60.775	13.880	16.762		20.09
1999	ILE	CG1	60.992	14.906	17.87		22.40
2000	ILE	CG2	59.660	14.316	15.820		17.07
2001	ILE	CD1	61.365	16.303	17.382		33.15
2002	LYS	N N	60.844	10.882	15.909		20.78
2003		CA	60.402	9.864 14.96		21.92	
2004		C	59.744	8.711 15.63		21.66	
2005		Ö	58.801	8.086 15.17		24.06	
2006		СВ	61.517	9.349 14.04		22.87	
2007		CG	62.275	10.451	13.370		20.42
2008	LYS	CD	63.483	9.834 12.67		33.57	
2009		CE	64:772	10.122			43.45
2010		NZ	65.169	9.089 14.39		74.10	
2011		N	60.328	8.382 16.73		19.22	
2012		CA	59.802	7.275 17.48		21.33	
2012		C	58.418	7.643 18.00		22.92	÷ •
2013	LYS	Ö	57.563	6.767 18.08		18.47	
2014		CB	60.804	6.907 18.59		26.85	
2016	LYS	CG	60.510	5.641 19.36		57.18	
2010	LIO		00.010	J.071 10.00	•	57.10	

	LYS	CD	61.693	5.220 20.236	74.44
2018	LYS	CE	61.283	4.289 21.393	100.00
2019	LYS	NZ	61.528	4.824 22.754	100.00
	VAL	N	58.209	8.943 18.386	20.32
2021	VAL	CA	56.880	9.344 18.869	19.06
2022		С	55.878	9.249 17.713	19.98
2023	VAL	0	54.733	8.767 17.821	19.01
2024	VAL.	CB	56.868	10.755 19.47	
2025		CG1	55.427	11.187 19.85	
2026	VAL	CG2	57.734	10.742 20.75	
2027	LEU	Ν	56.348		15.76
2028	LEU	CA	55.460		15.06
2029	LEU	С	55.086		21.46
2030	LEU	0	53.940		18.74
2031	LEU	CB	56.111	10.260 14.21	
2032	LEU	CG	55.289	10.130 12.91	
2033	LEU	CD1	53.915	10.764 13.09	
2034	LEU	CD2	56.004	10.808 11.71	1 17.04
2035	LEU	Ν	56.064	7.209 15.275	20.58
2036	LEU	CA	55.762	5.798 15.021	18.54
2037	LEU	Ç.	54.783	5.290 16.042	21.57
2038	LEU	0	53.935	4.472 15.713	22.28
2039	LEU	СВ	57.015	4.889 14.955	19.53
2040	ĿEU	CG	57.819	5.029 13.630	24.40
2041	LEU	CD1	59.230	4.498 13.813	22.15
2042	LEU	CD2	57.127	4.319 12.437	.21.17
2043	ASP	N	54.883	5.775 17.297	17.91
2044	ASP	CA	53.905	5.330 18.301	16.81
2045	ASP	С	52.538	5.845 17.897	19.40
2046	ASP	0	51.528	5.181 17.996	21.08
2047	ASP	CB	54.203	5.840 19.767	19.29
2048	ASP	CG	53.470	5.056 20.894	46.92
2049	ASP	OD1	53.626	3.832 21.196	39.76
2050	ASP	OD2	52.681	5.849 21.576	67.80
2051	MET	N	52.465	7.082 17.466	17.80
2052	MET	CA	51.146	7.550 17.093	18.08
2053	MET	С	50.586	6.785 15.902	19.16
2054	MET	0	49.360	6.544 15.802	19.67
2055	MET	CB	51.294	9.000 16.640	23.05
2056		CG	51.750	9.946 17.756	33.10
2057	MET	SD	51.452	11.712 17.33	8 37.23
2058	MET	CE	50.717	12.236 18.90	3 32.15
2059	ARG	N	51.471	6.425 14.953	16.98
2060	ARG	CA	50.996	5.690 13.759	18.78
	ARG	-C	50.448	4.309 14.082	20.60
2062	ARG	0	49.921	3.625 13.227	18.40
					•

2063	ARG	CB	51.817	5.867	12.441	25.17
			52.747	7.091	12.429	48.25
2064	ARG	CG				
2065	ARG	CD	52.842	8.056	11.215	60.15
2066	ARG	NE	52.920	7.361	9.965 30.10	
2067	ARG	CZ	53.076	7.701	8.652 25.89	
	ARG	NH1	53.321	8.890	8.086 27.29	
2068						
2069	ARG	NH2	53.031	6.649	7.819 19.72	47.55
2070	LYS	N	50.600	3.877	15.366	17.55
2071	LYS	CA	50.018	2.617	15.750	16.12
2072	LYS	,C	48.516	2.808	15.804	23.50
2073	LYS	0	47.753	1.864	15.744	21.94
2074	LYS	СВ	50.439	2.157	17.165	14.21
						18.36
2075	LYS	CG	51.927	1.901	17.250	
2076	LYS	CD -	52.363	1.502	18.666	21.09
2077	LYS	CE	53.863	1.184	18.762	28.07
2078	LYS	NZ	54.278	1.049	20.179	30.77
2079	PHE	N	48.066	4.046	15.996	17.28
2080	PHE	CA	46.649	4.276	16.181	17.04
2081	PHE	C	45.896	4.765	14.955	19.33
2082	PHE	Ö	44.665	4.622	14.894	17.66
				5.282	17.315	17.60
2083	PHE	CB	46.469			
2084	PHE	CG	47.096	4.756	18.576	17.54
2085	PHE	CD1	46.372	3.897	19.407	19.62
2086	PHE	CD2	48.368	5.176	18.950	18.66
2087	PHE	CE1	46.943	3.369	20.570	21.35
2088	PHE	CE2	48.910	4.698	20.145	22.41
2089	PHE	CZ	48.214	3.787	20.946	19.04
2090	ARG	N	46.598	5.382	14.053	13.88
	ARG	CA	45.967	5.775	12.828	13.48
				5.809	11.774	16.22
2092	ARG	C	47.051			
2093	ARG	0	48.163	6.250	11.973	15.25
2094	ARG	CB	45.321	7.172	12.939	16.93
2095	ARG	CG	44.544	7.632	11.689	16.86
2096	ARG	CD	43.697	8.897	12.048	18.74
2097	ARG	NE	42.861	9.407	10.923	16.22
2098	ARG	CZ	41.658		10.504	27.02
2099	ARG	NH1	41.047	7.860	11.071	15.47
		NH2	41.056		9.443 17.92	10.17
2100	ARG					18.09
2101	MET	N	46.715		10.564	10.09
2102	MET	CA	47.684		9.516 18.05	
2103	MET	C	47.968		8.978 20.84	
2104	MET	0	47.145	7.697	9.055 18.35	
2105	MET	СВ	47.079	4.556	8.343 19.02	
2106	MET	CG	45.812		7.720 21.47	
2107		SD	45.180	_	6.222 26.31	1 .1
	MET	CE	44.495		6.841 24.21	•
2108	IVIE I	UE	77.43J	2.102	U.UTI 24.21	

2109 2110 2111 2112 2113 2114 2115 2116	GLY GLY GLY GLY LEU LEU LEU	N CA C O N CA C O	49.125 49.540 49.743 49.791 49.925 50.072 51.314 52.374	6.961 8.363 8.083 7.529 9.433 8.135 10.451 9.400 9.463 10.648 11.339 10.727	18.21 21.75 7.402 18.48	14.70
2117 2118	LEU	CB CG	50.061 48.786		11.712 1 17.72 6 17.55	14.05
2119 2120 2121 2122 2123 2124	ILE ILE ILE ILE	CD1 CD2 N CA C	48.869 47.507 51.159 52.219 52.350 52.785	10.393 12.596 13.420 12.891 11.728	11.817 9.295 15.02 8.703 15.09 7.295 20.09 7.088 16.03	15.18
<ul><li>2125</li><li>2126</li><li>2127</li></ul>	ILE ILE ILE	CB CG1 CG2	53.538 53.244 54.476	13.451 14.065 14.391	9.423 17.27 10.772 8.650 17.72	16.88
2128	ILE GLN	CD1 N	54.453 51.917	14.681 13.710	11.446 6.331 18.36	18.43
2130 2131 2132	GLN GLN	CA C O	51.887 53.025 53.040	13.223 13.540 13.026	4.941 19.19 3.983 22.72 2.871 20.09	
2133 2134		CB CG	50.522 49.371	13.582 12.674	4.326 19.65 4.858 22.65	, Y) -
2135 2136 2137	GLN GLN GLN	CD OE1 NE2	49.367 49.184 49.600	11.349 11.364 10.236	4.113 26.84 2.900 17.52 4.801 19.94	
2138 2139	THR THR	N CA	53.911 55.045	14.414 14.841	4.397 21.73 3.612 20.72	·
<ul><li>2140</li><li>2141</li><li>2142</li></ul>	THR THR THR	C O CB	56.262 56.210 54.876	15.010 15.236 16.140	4.495 23.82 5.721 21.43 2.852 21.33	
2143 2144	THR THR	OG1 CG2	55.050 53.520	17.211 16.222	3.777 23.48 2.174 16.62 3.844 22.77	
<ul><li>2145</li><li>2146</li><li>2147</li></ul>	ALA ALA ALA	N CA C	57.378 58.601 58.801	14.910 15.018 16.403	4.587 23.01 5.105 22.21	
2148 2149 2150	ALA ALA ASP	O CB N	59.447 59.776 58.259	16.639 14.531 17.354	6.154 24.06 3.763 24.53 4.370 20.66	
2151 2152	ASP ASP	CA C	58.422 57.595	18.724 18.934	4.858 20.24 6.129 23.29	
2153 2154		O CB	57.899 58.049	19.740 19.778	7.024 23.21 3.806 23.48	

2155 2156 2157 2158 2159 2160 2161 2162 2163 2164 2165 2166	ASP ASP GLN GLN GLN GLN GLN GLN GLN GLN	CG OD1 OD2 N CA C O CB CG CD OE1 NE2	58.591 59.693 57.785 56.525 55.712 56.513 56.369 54.325 53.295 52.071 52.036 51.071	21.167 21.282 22.190 18.194 18.298 17.759 18.212 17.569 18.409 17.609 16.389 18.282	4.207 30.38 4.763 31.02 3.885 34.64 6.208 18.19 7.413 17.86 8.615 23.26 9.768 21.57 7.239 17.92 6.448 18.27 6.116 19.07 6.390 18.84 5.541 16.62	
2167 2168		N CA C	57.349 58.148 59.161	16.727 16.126 17.115	8.369 20.85 9.423 17.29 9.838 21.68	
2169 2170	LEU	0	59.368	17.386	11.015	21.08
2171	LEU	CB	58.770	14.818	8.982 16.92	10.01
2172		CG	59.679	14.141	10.018	18.94
2173		CD1	58.866	13.624	11.231 9.351 15.70	17.31
2174	LEU	CD2	60.409 59.781	12.959 17.705	8.820 20.27	
<ul><li>2175</li><li>2176</li></ul>	ARG ARG	N CA	60.792	18.727	9.101 19.02	
2177		CA	60.181	19.859	9.932 22.20	
2178	ARG	0	60.689	20.361	10.947	21.23
2179	ARG	CB	61.327	19.293	7.792 15.51	
2180	ARG	CG	62.568	20.163	8.046 21.78	
2181	ARG	CD	63.127	20.785	6.770 27.88	
2182	ARG	NE	64.351	21.523	7.038 33.02	
2183	ARG	CZ	64.355	22.814	7.227 31.22	
2184	ARG	NH1	63.242	23.517	7.186 23.06	
2185	ARG	NH2	65.508	23.412	7.478 31.35	
2186	PHE	N	59.007	20.291	9.471 20.79	
2187	PHE	CA	58.378	21.345	10.214	18.33
2188	PHE	C	58.180	20.945	11.680	23.47
2189	PHE	0	58.375	21.746	12.588	22.26
2190	PHE	CB	57.065	21.789	9.582 16.79 10.381	21.31
2191	PHE PHE	CG CD1	56.399 56.681	22.920 24.267	10.361	21.18
<ul><li>2192</li><li>2193</li></ul>	PHE	CD2	55.431	22.654	11.355	21.09
2193	PHE	CE1	56.063	25.309	10.814	22.40
2195	PHE	CE2	54.836	23.698	12.074	22.51
2196	PHE	CZ	55.164	25.025	11.833	19.57
2197	SER	N	57.752	19.714	11.884	20.57
2198	SER	CA	57.496	19.183	13.233	19.79
2199	SER	C	58.704	19.438	14.082	22.01
2200	SER	0	58.617	19.894	15.216	21.53

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2201	SER	СВ	57.252	17.666	13.231	16.35
2202	SER	OG	55.984	17.531	12.733	31.09
2203	TYR	Ν	59.841	19.086	13.544	20.69
2204	TYR	CA	61.030	19.291	14.365	19.51
2205	TYR	С	61.207	20.766	14.689	21.92
2206	TYR	0	61.557	21.130	15.818	20.31
2207	TYR	СВ	62.273	18.913	13.583	19.52
2208	TYR	CG	62.654	17.464	13.682	24.16
2209	TYR	CD1	61.877	16.472	13.074	24.17
2210	TYR	CD2	63.872	17.095	14.281	26.60
2211	TYR	CE1	62.245	15.126	13.119	24.68
2212	TYR	CE2	64.261	15.750	14.311	25.92
2213	TYR	CZ	63.440	14.771	13.749	24.97
2214	TYR	ОН	63.827	13.469	13.819	32.37
2215	ĿEU	N	60.985	21.623	13.670	19.51
2216	LEU	CA	61.150	23.020	13.916	17.52
2217	LEU	C	60.225	23.486	15.031	26.06
2218	LEU	Ō.	60.606	24.304	15.892	25.65
2219	LEU	СВ	60.860	23.865	12.721	16.93
2220	LEU	CG	61.898	23.718	11.592	26.47
2221	LEU	CD1	61.535	24.606	10.412	24.45
2222	LEU	CD2	63.302	24.080	12.029	28.00
2223	ALA	N	58.990	23.004	15.018	19.08
2224	ALA	CA	58.051	23.437	16.041	16.99
2225	ALA	С	58.424	22.901	17.408	20.57
2226	ALA	0	58.287	23.551	18.424	19.60
2227	ALA	СВ	56.587	23.121	15.708	16.41
2228	VAL	N	58.905	21.698	17.488	17.68
2229	VAL	CA	59.206	21.230	18.835	17.05
2230	VAL	C .	60.448	21.893	19.395	23.15
2231	VAL	0	60.592	22.155	20.616	21.24
2232	VAL	CB	59.378	19.694	18.826	20.45
2233	VAL	CG1	59.865	19.096	20.165	18.27
2234	VAL	CG2	58.084	19.006	18.357	19.14
2235	ILE	Ν	61.405	22.100	18.504	21.22
2236	ILE	CA	62.661	22.702	18.937	20.04
2237	ILE	С	62.420	24.090	19.475	25.88
2238	ILE	0	62.999	24.462	20.493	25.47
2239	ILE	CB	63.639	22.794	17.766	24.05
2240	ILE '	CG1	64.232	21.403	17.511	25.87
2241	ILE	CG2	64.736	23.758	18.113	22.69
2242	ILE	CD1	64.839	21.250	16.118	27.49
2243	GLU	N-	61.577	24.877	18.781	19.27
2244	GLU	CA	61.260	26.237	19.213_	17.92
2245	GLU	C	60.385	26.224	20.480	22.75
2246	GLU	0	60.556	27.020	21.388	20.84

2247	GLU .	СВ	60.510	26.936	18.054	19.09
2248	GLU	CG	59.998	28.359	18.389	20.36
2249	GLU	CD	61.125	29.327	18.744	29.49
2250	GLU	OE1	62.316	29.183	18.463	23.96
2251	GLU	OE2	60.693	30.344	19.428	26.29
2252	GLY	N	59.423	25.296	20.515	20.31
2253	GLY	CA	58.498	25.189	21.643	18.63
2254	GLY	C	59.259	24.833	22.907	23.95
2255	GLY	Ö	58.870	25.127	24.072	18.97
2256	ALA	N	60.356	24.110	22.677	23.58
2257	ALA	CA	61.188	23.677	23.804	23.19
2258	ALA	C	61.565	24.862	24.682	23.67
2259	ALA	Ö	61.732	24.708	25.900	25.14
2260	ALA	СВ	62.471	22.961	23.330	23.30
2261	LYS	N	61.747	26.038	24.042	19.35
	LYS	CA	62.155	27.232	24.809	19.49
2262			61.196	27.564	25.955	25.32
2263	LYS	С	61.575	27.937	27.090	22.02
2264	LYS	O	=	28.423	23.883	21.56
2265	LYS	CB	62.225		22.857	24.29
2266	LYS	CG	63.349	28.249	21.907	28.63
2267	LYS	CD	63.327	29.437	20.820	33.79
2268	LYS	CE	64.359	29.447		63.50
2269	LYS	NZ	63.978	30.470	19.839	
2270	PHE	N	59.922	27.482	25.608	20.00
2271	PHE	CA	58.858	27.740	26.571	17.26
2272		C	58.834	26.622	27.598	23.66
2273	PHE	0	58.789	26.847	28.791	23.24
2274	PHE	СВ	57.543	27.740	25.772	19.05
2275	PHE	CG	56.407	27.958	26.718	23.90
2276	PHE	CD1	55.800	26.859	27.315	30.06
2277	PHE	CD2	56.014	29.242	27.087	25.49
2278	PHE	CE1	54.788	27.010	28.262	31.06
2279	PHE	CE2	54.982	29.400	28.015	30.43
2280	PHE	CZ	54.380	28.294	28.612	28.57
2281	ILE	N	58.841	25.369	27.139	21.16
2282	ILE	ÇA .	58.815	24.235	28.070	21.71
2283	ILE	С	60.001	24.330	29.014	26.83
2284	ILE	0	59.949	23.959	30.185	28.05
2285	ILE	CB	59.014	22.890	27.335	24.04
2286	ILE	CG1	57.843	22.495	26.446	24.85
2287	ILE	CG2	59.269	21.758	28.309	26.71
2288	ILE	CD1	56.488	23.121	26.760	22.80
2289	MET	N	61.117	24.785	28.524	21.30
2290	MET	CA	62.205	24.825	29.440	23.12
2291	MET	C	62.325	26.098	30.299	28.91 <sup>,</sup>
2292	MET	0	63.398	26.410	30.748	31.20

2293	MET	СВ	63.520	24.378	28.836	25.89
2294	MET	CG	63.303	23.037	28.171	29.97
2295	MET	SD	63.168	21.686	29.352	29.79
2296	MET	CE	64.744	21.950	30.149	26.41
2297	GLY	N	61.278	26.868	30.528	24.26
2298	GLY	CA	61.489	27.964	31.450	24.71
2299	GLY	С	61.414	29.354	30.933	25.55
2300	GLY	0	61.441	30.290	31.708	24.08
2301	ASP	N	61.338	29.535	29.658	21.90
2302	ASP	CA	61.207	30.919	29.246	22.03
2303	ASP	С	59.786	31.173	28.742	27.04
2304	ASP	Ō	59.433	31.017	27.567	29.25
2305	ASP	СВ	62.264	31.272	28.189	25.34
2306	ASP	CG	62.034	32.645	27.603	36.54
2307	ASP	OD1	61.228	33.466	28.063	40.88
2308	ASP	OD2	62.728	32.823	26.512	33.74
2309	SER	N ·	58.906	31.542	29.605	24.08
2310	SER	CA	57.570	31.757	29.101	26.04
2311	SER	C	57.474	32.994	28.153	33.52
2312	SER	0	56.550	33.100	27.360	33.01
2313	SER	CB	56.523	31.840	30.246	28.08
2314	SER	OG	56.595	30.689	31.114	27.62
	SER	N	58.412	33.948	28.264	28.61
2315		CA	58.379	35.156	27.472	26.34
2316	SER				25.956	32.58
2317	SER	C	58.298	34.830		30.04
2318	SER	0	57.816	35.629	25.094	
2319	SER	CB	59.651	35.942	27.768	22.72
2320	SER	OG	60.712	35.599	26.864	34.10
2321	VAL	N	58.812	33.647	25.599	28.26
2322	VAL	CA	58.797	33.246	24.170	28.99
2323	VAL	C	57.356	33.239	23.563	32.46
2324	VAL	0	57.093	33.549	22.403	28.95
2325	VAL	СВ	59.628	31.965	23.969	29.56
2326	VAL	CG1	58.841	30.740	24.349	27.83
2327	VAL	CG2	60.017	31.836	22.548	31.39
2328	GLN	N	56.370	32.942	24.425	30.24
2329	GLN	.CA	54.974	32.874	24.017	29.27
2330	GLN	С	54.503	34.187	23.477	30.84
2331	GLN	0	53.777	34.236	22.506	27.71
2332	GLN	CB	54.073	32.406	25.207	32.06
2333	GLN	CG	52.561	32.481	24.958	56.39
2334	GLN	CD	51.805	31.561	25.885	51.02
2335	GLN	OE1	52.229	31.337	27.004	34.74
2336	GLN	NE2	50.703	31.010	25.403	62.41
2337	ASP	N	54.895	35.262	24.140	29.32
2338	ASP	CA	54.505	36.602	23.691	30.87

2339	ASP	С	55.136	36.911	22.333	30.01
2340	ASP	Ö,	54.530	37.499	21.433	31.24
2341	ASP	CB	54.906	37.629	24.756	35.79
2342	ASP	CG	54.358	37.219	26.082	65.63
2343	ASP	OD1	53.170	36.998	26.227	71.17
2344	ASP	OD2	55.278	37.034	27.014	80.27
2345	GLN	N	56.374	36.460	22.194	25.78
2346	GLN	CA	57.108	36.574	20.948	26.06
2347	GLN	C	56.384	35.851	19.806	30.03
2348	GLN	0	56.246	36.357	18.715	26.89
2349	GLN	CB	58.455	35.924	21.120	27.67
2350	GLN	CG	59.317	36.611	22.198	44.36
	GLN	CD	60.751	36.116	22.147	77.73
2351		OE1	61.283	35.846	21.053	88.38
2352	GLN			35.840 35.952	23.326	61.19
2353	GLN	NE2	61.357 55.906	34.644	20.070	26.22
2354	TRP	N	55.195		19.048	23.95
2355	TRP	CA		33.953	19.046 18.716	23.93 27.74
2356	TRP	C	53.959	34.709		
2357	TRP	0	53.590	34.825	17.557	29.34
2358	TRP	CB	54.752	32.529	19.487	22.50
2359	TRP	CG	55.887	31.639	19.847	21.30
2360	TRP	CD1	57.167	31.756	19.417	23.75
2361	TRP	CD2	55.842	30.526	20.763	19.48
2362	TRP	NE1	57.927	30.767	19.994	23.42
2363	TRP	CE2	57.143	30.005	20.849	22.89
2364	TRP	CE3	54.820	29.939	21.517	20.00
2365	TRP	CZ2	57.464	28.895	21.668	20.62
2366	TRP	CZ3	55.119	28.829	22.299	20.58
2367	TRP	CH2	56.418	28.303	22.365	20.81
2368	LYS	N	53.273	35.191	19.732	25.02
2369	LYS	CA	52.051	35.913	19.449	23.72
2370	LYS	С	52.279	37.141	18.512	32.33
2371	LYS	0	51.536	37.430	17.531	28.38
2372	LYS	CB	51.390	36.292	20.731	24.86
2373	LYS	CG	50.049	36.954	20.489	36.15
2374	LYS	CD	49.612	37.791	21.672	56.07
2375	LYS	CE	48.105	37.986	21.748	94.91
2376	LYS	NZ	47.664	38.697	22.968	100.00
2377	GLU	N	53.319	37.895	18.808	32.11
2378	GLU	CA	53.630	39.036	17.969	35.38
2379	GLU	С	54.028	38.602	16.559	36.31
2380	GLU	0	53.557	39.097	15.540	38.26
2381	GLU	CB	54.813	39.855	18.548	39.32
2382	GLU	CG	54.559	40.416	19.974	69.72
2383	GLU	CD	53.542	41.540	19.993	100.00
2384	GLU	OE1	53.826	42.742	19.907	100.00

2385	GĽU	OE2	52.311	41.071		20.08	6	100.00
2386	LEU	N	54.930	37.668		16.47		29.59
2387	LEU	CA	55.371	37.218		15.17		29.45
2388	LEU	C	54.297	36.602		14.31		34.02
				36.402		13.09		34.53
2389	LEU	0	54.438			15.38		30.02
2390	LEU	CB	56.402	36.124				37.18
2391	LEU	CG	57.673	36.730		15.95		
2392	LEU	CD1	58.611	35.621		16.44		36.22
2393	LEU	CD2	58.332	37.541		14.83		39.33
2394	SER	N	53.238	36.196		14.95		26.99
2395	SER	CA	52.226	35.561		14.17		27.01
2396	SER	С	51.217	36.583		13.60		33.85
2397	SER	0	50.359	36.208		12.84	3	33.74
2398	SER	CB	51.521	34.632		15.13	4 -	28.99
2399	SER	OG	50.444	35.407		15.64	8	44.35
2400	HIS	Ν	51.275	37.855		14.03	4	33.50
2401	HIS	CA	50.355	38.935		13.60	4	36.33
2402	HIS	C	48.945	38.510		13.66		34.53
2403	HIS	Ö	48.221	38.641		12.66		29.05
2404	HIS	CB	50.574	39.458		12.17		41.53
2405	HIS.	CG	52.003	39.597	•	11.96		50.56
2406	HIS	ND1	52.683	38.801		11:04		55.72
	HIS	CD2	52.889	40.377		12.65		55.69
2407				39.152		11.15		56.62
2408	HIS	CE1	53.978			12.12		56.78
2409	HIS	NE2	54.131	40.085				31.47
2410	GLU	N	48.528	38.023		14.79		
2411	GLU	CA	47.172	37.560		14.77		32.57
2412	GLU	C	46.110	38.585		14.65		35.60
2413	GLU	0	44.984	38.253		14.39		35.07
2414	GLU	CB	46.886	36.568		15.88		34.05
2415	GLU	CG	47.343	37.124		17.21		29.10
2416	GLU	CD	46.766	36.272		18.29	1	44.34
2417	GLU	OE1	46.803	35.044		18.31	1	28.10
2418	GLU	OE2	46.156	36.987		19.18	3	42.70
2419	ASP	N <sup>ii</sup>	46.433	39.851	•	14.84	1	38.45
2420	ASP	CA	45.375	40.870		14.77	7	78.44
2421	ASP	С	45.114	41.525		13.42	8	73.26
2422		0	46.027	41.652		12.62	0	57.37
2423		СВ	45.440	41.851		15.94	9	80.14
2424		CG	45.045	41.139		17.21		99.39
2425	ASP	OD1	43.981	40.514	1.	17.33		100.00
2426		OD2	45.979	41.210		18.14		100.00
2427	HOH	.0	38.401	17.896		24.63		19.06
2427	НОН	0	42.880	-2.994 ·			22.22	10.00
	пОп .НОН		- 37.909 -	 0.024			18.06	
	HOH.	0	34.283	 3.652			16.00	
2430	пОП	J	J4.Z0J	3.032	10.00	I.	10.33	

2431 2432 2433 2434 2435	HOH HOH HOH HOH	0 0 0 0	31.031 56.762 38.520 38.706 48.541	7.792 20.003 16.414 4.591 3.340 8.875 4.448 17.369	35.819 17.34	23.22
2436	HOH	0	22.375	24.091 3.090 10.757	20.022	27.04
2437	HOH	0	50.383		, 19.44 34.028	23.47
2438	HOH	0	56.581	20.545	27.502	26.90
2439	HOH	0	44.023	22.433	15.041	26.35
2440	HOH	0	23.678	29.248		25.48
2441		0	30.955	16.693	24.220 9         27.62	
2442	HOH	0	36.539	-2.273 15.479	6.928 25.25	
2443	HOH	0	60.199		24.284	24.08
2444	HOH	0	42.799	15.905	13.755	
2445	HOH	0	58.854	31.126	9.898 29.35	
2446	HOH	0	25.489	12.276	30.537	27.95
2447		0	44.868	21.909	16.536	
2448	HOH	0	19.794	30.078 3.661 14.164		
2449	HOH	0	41.421	2.616 3.383		,
2450	HOH	0	41.884 34.858	3.969 20.688		
2451	HOH	0		2.425 13.962		
2452		0	53.734	9.816 23.103		
2453	HOH	0	60.930	3.496 20.674		
2454	HOH	0	42.886			26.91
2455	HOH	0	30.810		18.288	26:16
2456	HOH	0	59.810	32.233	32.172 1.262 32.83	
2457	HOH	0	57.738	17.405	5.425 23.54	
2458	HOH	0	36.660	27.108	1.767 24.56	-
2459	HOH	0	32.265	17.633	2.407 38.58	
2460	HOH	0	66.198	17.107	31.074	29.94
2461	HOH	0	57.752	28.197 15.987	26.904	32.37
2462	HOH	0	41.286	27.749	2.270 31.44	
2463	HOH	0	36.669 39.624	-4.625 16.50		
2464 2465	HOH HOH	0	45.398	24.023	0.668 27.71	
2466	HOH	0	51.673	32.445	21.771	23.43
2467	HOH	0	53.099	24.833	2.471 29.74	
2468	HOH	0	54.526	19.450	2.810 28.76	
2469	HOH	Ö	27.105	36.172	19.570	24.82
2470	HOH	Ö	36.334	30.259	9.265 30.82	
2471	HOH	Ö	56.626	24.387	33.815	30.17
2472	HOH	0 .	42.738	29.954	24.494	35.73
2473	HOH	0	57.220	13.971	0.977 32.27	
2474	НОН	Ö	65.256	29.469	30.089	39.20
2475		0	49.786	39.709	16.909	43.80
2476	НОН	Ö	52.863	6.253 24.788		
0		<del>-</del>				





2477	HOH	0	35.798	-5.535 5.469	31.13	
2478	НОН	0	50.331	22.923	-0.657 36.76	
2479	НОН	0	36.765	3.406 1.739	42.22	
2480	НОН	Ö	26.434	4.890 5.483	35.83	
2481	НОН	Ŏ	34.613	32.955		43.11
2482	НОН	Ö	27.837	3.167 1.403		
2483	HOH	0 -	32.582	9.372 -0.315		
	НОН	0	45.800	3.147 2.702		
2484	НОН			0.985 2.765		
2485		0		1.300 20.11		
2486	HOH	0 .	29.040	-4.2226.234		
2487	HOH	0	42.199			
2488	HOH	0	21.619	20.058	9.321 40.28	
2489	НОН	0	53.500	20.656	27.642	27.42
2490	HOH	0	16.860	27.164	18.394	
2491	HOH	0	43.874		3 25.50	
2492	HOH	0	29.683	-0.569 10.39		
2493	HOH	0	28.054	11.908	26.787	41.40
2494	HOH	0	50.466		28.067	42.56
2495	HOH	0	28.502	21.053	3.156 30.33	
2496	HOH	0	63.942	27.419	17.604	28.20
2497	HOH	0	22.109	21.232	30.110	48.48
2498	HOH	0	49.254	32.003	23.342	40.08
2499	HOH	0	24.641	7.692 4.830	53.24	
2500	HOH	Ò	63.797	11.089	16.405	27.62
2501	HOH	0	53.333	9.033 24.88	1 28.87	
2502	HOH	0	37.700	0.242 19.12	7 49.32	
2503		0	24.665	29.142	9.720 48.36	
2504	НОН	0	54.352	-3.1149.194	44.58	
2505	HOH	Ö	62.631	5.916 11.92		
	НОН		63.952	28.080	27.641	38.19
2507	НОН		65.802	19.783	33.411	38.59
		Õ	57.255		7.852 41.05	
	НОН	•	19.030	25.560	24.767	57.85
2510	HOH		64.201		18.348	29.38
2511	HOH	Ö	55.852	-0.42722.99		
2512		0	36.898	-5.489 16.05		•
2512	HOH	0	33.905	27.753	6.095 27.01	
				-1.372 13.74		
2514	HOH	0	27.382	·	8 41.00	
2515	HOH	0	33.489			
2516	HOH	0	24.559	2.258 7.150		25 04
2517	HOH	0	65.779	15.180	21.392	35.84
2518	HOH	0	65.553	25.125	21.639	47.37
2519	HOH	0	32.513	19.611	26.519	62.17
2520	HOH	0	33.651	17.294	28.933	55.49
2521	HOH		41.137	19.611	-7.004 52.94	
2522	HOH	0	66.395	21.128	4.224 47.91	

2523	НОН	0	23.251	11.180	19.061	48.42
2524	HOH	0	39.259	33.229	22.025	51.45
2525	HOH	0 .	36.119	32.908	9.928 44.34	
2526	НОН	0	53.725	36.111	5.937 100.00	
2527	HOH	0	63.179	35.153	24.999	55.42
2528	HOH	0	35.968	6.586 -0.582		
2529	HOH	0	21.346		56.07	
2530	HOH	0 -	19.175		19.382	53.38
2531	HOH	0	22.715		5 50.52	
2532	HOH -	0	47.395	34.116		47.24
2533	HOH	0	32.765		3 64.04	
2534	HOH	0	54.471	34.005	11.028	47.55
2535	HOH	0	72.465	12,809	15.523	46.13
2536	HOH	0	47.441	6.136 -1.870		
2537	HOH	0	43.416	18.301	-3.45649.44	
2538	HOH	0	65.579	21.420	21.981	50.95
2539	HOH	0	47.751	7.411 2.407	40.84	
2540	HOH	0	32.644	8.712 8.756	15.74	•
2541	HOH	0	33.023	11.570	8.792 15.03	
2542	НОН	0 .	44.089	25.927	7.019 19.82	
2543	НОН	0	58.775	26.249	32.830	26.89
2544	HOH	0	51.112	4.543 8.642	20.55	
2545		0	62.946	25.839	15.605	26.99
2546	НОН	0	40.881	-6.288 14.710	the state of the s	
2547	НОН	0	52.351	8.308 5.018	25.66	
2548	НОН	0	61.224	24.419		30.44
2549	НОН	0	28.243	-3.9094.213		
2550	НОН	Ō.	62.221	27.938	13.810	30.53
2551	НОН	O	39.610	-6.610 12.43	1 30.17	
2552	НОН	Ö	54.043	16.770	35.701	34.80
2553	НОН	Ö	15.219		20.359	31.31
2554	НОН	Ö	64.424	20.767	35.354	32.43
2555	НОН	Ö	14.987	23.468	24.936	33.81
2556	НОН	0	21.319	29.054	23.219	34.70
2557	НОН	Ö	30.027	-1.696 12.465		• •
2558	НОН	Ö	25.366	15.033	7.774 33.88	
2559	НОН	Ö	66.346	11.979	1.208 42.20	
2560	HOH	Ö	17.665	23.698	26.769	54.87
2561	HOH	Ö	51.859	5.861 4.981		,
2562	HOH	Ö	24.749	29.791	25.560	42.21
2563	НОН	0	57.813	3.867 18.81		72.2
2564	HOH	0	63.315	26.715	7.503 46.17	
	HOH	0	53.229	27.524	4.226 50.52	
2565	НОН			7.825 9.883		
2566			63.148		7.236 22.65	de vinn 1.
2567		0	26.879	19.362	34.499	44.34
2568	HOH	0	67.878	17.615	34.433	44.34

2569 2570 2571	HOH HOH HOH	0 0 0	54.054 54.931 44.558		31.561 0.470 42.81 370.17	
2572		Ŏ	26.495		25.981	49.19
2573		Ŏ	45.561		6 47.40	
2574		Ŏ	33.903			
	НОН	Ö .	64.772	6.818 21.874	4 47.22	
2576		0	38.599	3.739 20.598	47.72	
2577	HOH	0	40.578	33.268	10.218	43.71
2578	HOH	0	23.004	14.669	9.548 41.06	
2579	HOH	0	18.520	21.210	10.100	82.04
2580	HOH	0	54.443			
2581	HOH	0	53.301	3.094 5.341	46.46 55.82	
2582	HOH	0	62.577	30.234	15.315	
2583	HOH	0	58.940		5.711 51.05	
2584	HOH	0	20.726	3.366 15.60		
	HOH	0	43.027	13.430	27.118	54.39
2586		0	62.195	28.464	10.795	46.23
2587		0	50.510		10.824	
	НОН	0	40.918		9 65.28	
2589	НОН	0			-2.42847.56	50.00
2590	HOH	0	13.754	15.204		53.86
2591		0	31.164		3.629 56.80	
2592		0	64.794	14.809	1.227 46.20	
2593	HOH	0 .	••••	6.902 9.576		
2594	HOH	0	22.779	9.377 7.806		
2595	HOH	0	35.536		2 58.11 14.904	65.02
2596	HOH	0	48.720	41.298	14.904	
2597	HOH	0	41.886		8.779 67.26	
2598 2599	HOH HOH	0	21.553 47.438		-1.762 68.04	
2600	НОН	0	59.462		2 44.01	
2601	HOH	0	34.650	13.050	10.584	15.74
2602	HOH	0	44.151	3.876 10.554		10.7
2603	НОН	0	30.226	8.145 9.680	•	
2604	HOH	Ö	50.795	5.197 23.429		
2605	НОН	Ö	55.241		1.478 43.03	
2606	НОН	Ö	13.876	26.053	19.951	39.87
2607	НОН	Ö	37.440	7.794 22.99		
2608	НОН	0	36.776	4.942 22.48		
2609	НОН	Ô	49.204	6.833 4.042		
2610	HOH	Ο.	46.016	15.869°	28.666	53.22
2611		0	52.597	14.225	35.410	44.55
2612	HOH	0	55.717	8.332 23.77	0 40.53	
2613	HOH	0	35.617	00.011	15.729	45.63
2614	HOH	0	24.347	36.908	13.597	48.19



2615	НОН	0	37.734	23.590	30.260	62.04
2616	HOH	0	59.359	7.814 23.35	1 51.17	•
2617	HOH	0	40.870	-8.615 10.30	1 39.22	) •
2618	HOH	0	55.653	-1.953 5.995	52.57	
2619	HOH	0	51.582	1.831 22.83	8 43.21	
2620	HOH	0	23.536	16.687	3.287 66.26	i
2621	НОН	0	27.390	9.583 26.22	2 66.80	)
2622	НОН	0	26.006		7.979 55.20	1
2623	НОН	O.	48.629	36.281	9.258 53.85	,
2624	HOH	0	54.312	1.731 24.31	2 49.12	
2625	НОН	0	64.534	7.113 16.40	7 42.12	
2626	НОН	Ō	43.617	30.982	3.143 48.34	
2627	НОН	Ō	22.572	13.132	27.363	70.18
2628	НОН	Ö	70.408	21.965	2.674 89.51	
2629	НОН	Ö	63.931	33.405	22.517	55.19
2630	НОН	Ö	48.600		29.774	71.55
2631	НОН	Ŏ	34.250	14.917	30.699	46.61
2632	НОН	Ö	43.731	33.337	6.158 93.58	
2633	НОН	Ö	60.489	31.207	10.548	EE 40
2634	НОН		50.155	2.150 6.615		
2635	НОН	0	25.514		2 75.79	ı
2636	НОН	Ö	31.745	30.209	9.802 50.27	
2637	НОН	Ö	35.679	10.169	26.838	65.87
2638	HOH	0	13.764	16.372	28.336	78.20
2639	HOH		17.554	19.418	30.248	64.92
2640	HOH	0	65.875	16.089	34.874	64.83
2641	HOH	0,	68.340		27.806	60.04
2642	HOH	0	36.275	32.819	22.710	48.84
2643	HOH	0	63.468		8 51.50	
2644	HOH	0	39.850	34.128	24.551	67.42
2645	HOH	0	22.019		7.060 70.82	
2646	HOH	0	30.890	-4.164 13.15		
		0	23.362	5.115 3.428		
2647	HOH		41.079	12.774	28.276	77.64
2648	HOH .			37.768	15.682	60.54
2649		0	32.157	12.911	33.109	69.06
2650	HOH	0	37.346		,	
2651	HOH	0	60.400	-0.560 17.55	1.856 20.00	
1	95 05	C1	45.324	12.023	3.269 20.00	
2	95	S2	45.841	12.927	2.692 20.00	
3	95	C3	44.986	14.367		
4	95	C4	44.391	14.159	1.498 20.00	
5	95	C5	44.709	12.824	0.981 20.00	
6	95 .	N6	44.964		3.482 20.00	
7	95	C7	45.489	15.733	4.720 20.00	
8	95	-C8	45.238	16.996	5.547 20.00	
9	95	09	44.228	17.659	5.375 20.00	

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10	95	O10	45.978	17.336	6.420 20.00
11	95	011	46.159	14.860	5.251 20.00
12	95	C12	43.503	15.119	0.776 20.00
13	95	O13	43.428	15.102	-0.452 20.00
14	95	014	42.803	15.898	1.358 20.00
15	95	C18	44.232	12.358	-0.368 20.00
16	95	C19	44.430	10.836	-0.521 20.00
17	95	N20	45.675	10.345	0.087 20.00
18	95	C21	45.756	10.616	1.531 20.00



Table of the orthogonal three dimensional coordinates in Ångstroms and B factors (Ų) for Protein Tyrosine Phosphatase 1B complexed with 7-(5-methoxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (Example 26).

No	Amin	o acid	X	Υ	Z	В			
1	GLU	N	39.465		52.977		54.384		50.84
2	GLU	CA	38.798		51.719		54.051		51.38
3	GLU	C	39.109		51.267		52.590		49.72
4	GLU	0	38.944		52.002		51.620		49.04
5	GLU	CB	37.270		51.855		54.252		53.79
6	GLU	CG	36.557		50.495		54.416		57.54
7	GLU	CD	36.972		49.698		55.707		62.60
8	GLU	OE1	36.770		50.216		56.799		64.63
9 .	GLU	OE2	37.518		48.585		55.616		64.40
10	MET	N	39.495		49.971		52.496		46.10
11	MET	CA	39.753		49.178		51.264		42.61
12	MET	С	38.547		49.091		50.363		40.49
13	MET	0	38.640		48.979		49.159 ·		38.08
14	MET	CB	40.128		47.730		51.640		41.62
15	MET	CG.	39.190		47.107		52.716		40.45
16	MET	SD -	39.645		45.471		53.299		35.84
17	MET	CE	41.295	•	45.867	•	53.958		39.51
18	GLU	N	37.419		49.189		51.051		40.04
19	GLU	CA	36.086		49.198		50.457		41.99
20	GLU	С	35.767		50.510		49.639		41.64
21	GLU	0	35.409		50.486		48.472		42.33
22	GLU	CB	35.156		48.876		51.627		43.82
23	GLU	CG	33.862		48.113		51.292		47.17
24	GLU	CD	33.244		47.563		52.607	•	49.31
25	GLU	OE1	33.732		47.923		53.672		48.56
26	GLU	OE2	32.301		46.761		52.564		51.19
27	LYS	Ν	36.000		51.696		50.213		40.99
28	LYS	CA	35.753		52.839		49.308	•	41.50
29	LYS	С	36.854		52.957		48.192		39.57
30	LYS	0	36.534		53.231		47.054		39.12
31	LYS	CB	35.643		54.142	•	50.104	. •	45.89
32	LYS	CG	34.578		54.185		51.248		51.71
33	LYS	CD	35.008		55.173		52.386		56.13
34	LYS	CE	34.450		54.867		53.793		59.81
35	LYS	NZ	35.323		55.409		54.876		62.35
36	GLU	N	38.147		52.704		48.526		38.42

37	GLU	CA	39.236	52.572	47.517	37.30
38	GLU'	C	38.888	51.620	46.385	34.58
39	GLU	0	39.054	51.964	45.238	33.74
		CB	40.601	52.140	48.101	40.63
40	GLU		•		47.019	46.16
41	GLU	CG	41.526	51.508		
42	GLU	CD	43.077	51.484	47.114	49.22
43	GLU	OE1	43.672	50.792	47.941	50.86
44	GLU	OE2	43.723	52.146	46.288	51.79
45	PHE	N	38.395	50.433	46.755	31.46
46	PHE	CA	37.835	49.488	45.799	30.49
47	PHE	С	36.777	50.107	44.857	33.05
48	PHE	0	36.828	49.954	43.648	31.17
49	PHE	CB	37.148	48.330	46.531	26.00
50	PHE	CG	36.682	47.268	45.560	23.39
51	PHE	CD1	37.608	46.321	45.069	24.75
52	PHE	CD2	35.388	47.202	45.084	23.83
53	PHE	CE1	37.277	45.364	44.109	21.69
54	PHE	CE2	35.064	46.263	44.099	23.15
55	PHE	CZ	35.985	45.348	43.603	22.50
56	GLU	N	35.818	50.823	45.452	34.78
57	GLU	CA	34.725	51.349	44.619	36.92
58	GLU	C	35.092	52.587	43.702	34.88
59	GLU	Ö	34.771	52.673	42.529	34.89
60	GLU	СB	33.592	51.542	45.609	42.69
61	GLU	CG	33.191	50.163	46.129	51.81
62	GLU	CD	32.044	50.199	47.121	58.22
63	GLU	OE1	31.735	51.274	47.661	60.76
64	GLU	OE2	31.467	49.123	47.337	60.88
65	GLN	N	35.875	53.523	44.252	32.17
66	GLN	CA	36.614	54.515	43.493	32.79
67	GLN	C	37.270	53.878	42.281	32.59
68	GLN	Ö	37.068	54.411	41.198	33.38
69	GLN	СВ	37.684	55.103	44.398	38.01
70	GLN	CG	38.460	56.386	44.041	46.72
71	GLN	CD	39.538	56.621	45.198	54.60
				56.281	45.103	58.48
72	GLN	OE1	40.705		46.335	57.65
73	GLN	NE2	39.104	57.129 52.753	42.495	30.75
74	ILE	N	37.995	52.752		29.79
75	ILE	CA	38.784	52.110	41.427	
76	ILE	C	37.944	51.513	40.345	29.61
77	ILE	0	38.222	51.644	39.164	28.92 28.01
.78	ILE	CB	39.820	51.020	41.780	
79	ILE	CG1	40.944	51.578	42.625	26.65
80	ILE	CG2	40.547	50.559	40.485	24.67
81	ILE	CD1	41.717	50.482	43.321	30.61
82	ASP	N	36.941	50.841	40.826	30.56

83	ASP	CA	36.056	5	0.131	•	39.941	•	34.30
84	ASP	C	35.275		1,174		39.066		36.61
85	ASP	Ō	35.220		1.032	3	37.847		37.72
86	ASP	СВ	35.183		9.189	4	10.820		34.23
87	ASP	CG	35.372		7.692	4	40.610		34.09
88	ASP	OD1	36.406		7.266	4	40.156		34.45
89	ASP	OD2	34.468		6.917		40.892		35.24
90	LYS	N	34.750		2.271		39.708		38.57
91	LYS	CA	34.018		3.296		38.901	•	40.78
92	LYS	C	34.892		3.932		37.806		40.89
93	LYS	0	34.499		4.071		36.665		41.19
94	LYS	·CB·	33.321		4.430		39.687		44.73
95	LYS	CG	34.160		5.716		39.765		49.46
96	LYS	CD	33.790		6.625		10.936		52.61
97	LYS	CE	34.890		7.651		11.277		51.86
98	LYS	NZ	36.251		7.064		41.269		49.92
99	SER	N .	36.087		4.345		38.162		39.93
100	SER	CA	36.820		4.789		37.003		41.11
101	SER	C <sub>.</sub>	37.425		3.593		36.217		41.60
101	SER	0	37.968		3.748		35.149		44.13
103	SER	СВ	37.986		5.539		37.561		41.13
103	SER	OG	38.877		4.652		38.356		43.06
105	GLY	N	37.399		2.386		36.767		39.95
106	GLY	CA	38.129		1.274		36.127		36.83
107	GLY	C	39.671		1.290		36.223		35.39
108	GLY	Ö	40.316		0.783		35.320		37.15
109	SER	Ň	40.279		1.841		37.301		32.66
110	SER	CA	41.780		1.938		37.308		31.67
111	SER	C	42.598		0.665		37.811		29.26
112	SER	Ö	43.741		0.778		38.218		28.04
113	SER	СB	42.337		3.166	(	38.097		32.05
114	SER	ŌĞ	41.454		4.345		38.032		36.33
115	TRP	N	42.019	4	9.485		37.773		26.65
116	TRP	CA	42.787	4	8.357		38.207		21.95
117	TRP	С	44.128	4	8.196		37.429		21.43
118	TRP	0	45.199	4	8.050		38.019		21.86
119	TRP	CB	41.882	4	7.101	(	38.149		22.70
120	TRP	CG	40.932	4	7.148		39.318		20.91
121	TRP	CD1	39,592	4	7.426	•	39.181	•	21.20
122	TRP	CD2	41.184	. 4	7.089	.4	40.757		21.02
123	TRP	NE1	39.031	4	7.559		40.408		21.97
124	TRP	CE2	39.970	4	7.310		41.388	. *	20.57
125	TRP	CE3	42.283		6.789		41.551	•	18.68
126	TRP	CZ2	39.847		7.294		42.770		20.86
127	TRP	CZ3	42.185		6.754		42.958		16.67
128	TRP	CH2	40.956	4	6.981	4	<b>43.559</b> .		19.85

129	ALA	N	44.103	48.273	36.096	21.75
130	ALA	CA	45.413	48.072	35.405	21.29
131	ALA	C	46.515	49.117	35.696	19.48
132	ALA	Ō	47.664	48.776	35.736	17.73
133	ALA	СВ	45.192	47.958	33.915	21.57
134	ALA	N	46.103	50.326	35.925	21.34
135	ALA	CA	46.952	51.428	36.319	20.36
136	ALA	C	47.471	51.300	37.816	20.18
137	ALA	Ö	48.633	51.406	38.160	23.47
138	ALA	СB	45.932	52.549	36.136	21.69
139	ILE	N .	46.537	50.973	38.753	20.36
140	ILE	CA	47.001	50.649	40.115	20.21
141	ILE	C	48.074	49.511	40.105	19.21
142	ILE	Ö	49.155	49.583	40.676	20.20
143	ILE	СВ	45.776	50.220	40.970	21.58
144	ILE	CG1	44.697	51.339	41.220	25.33
145	ILE	CG2	46.173	49.573	42.320	23.50
146	ILE	CD1	45.051	52.433	42.267	26.28
147	TYR	N	47.688	48.424	39.421	18.90
148	TYR	CA	48.574	47.264	39.255	16.59
149	TYR	C	49.878	47.625	38.498	19.69
	TYR	0	50.967	47.421	38.997	21.05
150		СВ	47.748	46.176	38.592	15.48
151	TYR		48.537	44.932	38.537	15.58
152	TYR	CG CD1	48.907	44.332	39.665	15.71
153	TYR	CD1	48.960	44.470	37.316	17.07
154	TYR	CE1	49.695	43.057	39.598	15.33
155	TYR	CE2	49.775	43.341	37.218	18.24
156	TYR TYR	CZ	50.173	42.628	38.344	17.83
157		OH	50.173	42.020	38.105	16.43
158	TYR GLN	N	49.759	48.251	37.330	20.71
159	_		50.882	48.930	36.627	23.54
160	GLN	CA		49.705	37.616	21.33
161	GLN	С	51.839	49.705 49.495	37.568	22.52
162	GLN	0	53.047	49.495 49.816	35.499	29.74
163	GLN	CB	50.268		34.238	43.77
164	GLN	CG	49.671	49.057	34.236 33.267	50.76
165	GLN	CD	48.539	49.650	33.495	53.37
166	GLN	OE1	47.671	50.472	32.066	51.41
167	GLN	NE2	48.508	49.076	38.587	22.69
168	ASP	N	51.244	50.451	39.649	23.75
169	ASP	CA	51.903	51.194		21.78
170	ASP	С	52.715	50.305	40.609 40.718	21.76
171	ASP	O	53.932	50.387	40.716	26.73
172	ASP	CB	50.890	52.087	39.655	31.02
173	ASP	CG OD1	50.646	53.429	39.035 39.025	31.02
174	ASP	OD1	51.574	53.975	33.023	31.00

175	ASP	OD2	49.544	53.975	39.841	32.48
176	ILE	Ν	52.020	49.333	41.202	20.63
177	ILE	CA	52.741	48.290	41.962	19.38
178	ILE	С	53.874	47.578	41.116	19.66
179	ILE	0	54.989	47.370	41,576	18.07
180	ILE	CB	51.688	47.279	42.579	19.75
181	ILE	CG1	50.762	48.016	43.509	19.61
182	ILE	CG2	52.236	46.021	43.304	16.65
183	ILE	CD1	49.363	47.437	43.437	21.44
184	ARG	Ν	53.565	47.169	39.884	19.64
185	ARG	CA	54.567	46.484	39.085	22.20
186	ARG	С	55.816	47.437	39.067	22.74
187	ARG	0	56.932	46.979	39.099	20.20
188	ARG	CB	54.023	46.237	37.637	23.80
189	ARG	CG	53.320	44.918	37.290	29.28
190	ARG	CD	53.204	44.519	35.775	38.17
191	ARG	NE	52.689	43.130	35.686	47.71
192	ARG	CZ	53.098	42.163	34.783	46.22
193	ARG	NH1	53.704	42.464	33.629	51.10
194	ARG	NH2	52.965	40.881	35.132	39.82
195	HIS	N	55.615	48.784	39.064	24.89
196	HIS	CA	56.730	49.740	38.862	28.65
197	HIS	C	57.493	50.276	40.108	27.86
198	HIS	Ö	58.673	50.654	40.038	29.52
	HIS	СВ	56.179	50.983	38.193	36.09
200	HIS	CG	56.811	51.114	36.834	47.89
201	HIS	ND1	58.149	50.961	36.592	52.16
202	HIS	CD2	56.145	51.425	35.625	50.80
203	HIS	CE1	58.303	51.179	35.262	54.85
204	HIS	NE2	57.113	51.465	34.656	55.16
205	GLU	N	56.716	50.341	41.204	24.60
206	GLU	CA	57.222	50.671	42.538	23.35
207	GLU	C	57.907	49.520	43.236	20.84
208	GLU	Ö	58.770	49.755	44.059	20.77
209	GLU	СВ	56.078	51.097	43.441	25.78
210	GLU	CG	55.482	52.418	43.015	34.71
211	GLU	CD	54.294	52.679	43.916	41.97
212	GLU	OE1	54.457	52.660	45.153	47.08
213	GLU	OE2	53.218	52.905	43.375	42.03
214	ALA	N	57.458	48.299	42.919	20.22
215	ALA	CA	58.071	47.145	43.572	18.85
216	ALA	C	59.618	47.108	43.413	19.60
217	ALA	Ö	60.137	47.773	42.539	21.31
218	ALA	СВ	57.514	45.879	42.928	17.65
219	-SER	-N	60.263	46.351	44.323	17.44
220	SER	CA	61.700	46.463	44.627	17.65
220	OLIV	<b>U</b> / (	31.700	10,700		

221	SER	С	62.475	45.590	43.693	20.44
222	SER	0	61.875	44.737	43.028	20.82
223	SER	CB	61.987	45.892	46.086	16.87
224	SER	OG	60.908	46.087	47.088	17.03
225	ASP	N	63.806	45.769	43.757	23.65
226	ASP	CA	64.608	44.993	42.881	25.32
227	ASP	C	65.953	44.653	43.545	24.00
228	ASP	Ö	66.834	45.495	43.638	25.47
229	ASP	ĊB	64.577	45.897	41.647	30.03
230	ASP	CG	65.409	45.356	40.505	35.94
231	ASP	OD1	65.782	44.168	40.576	37.82
232	ASP	OD1	65.652	46.122	39.542	40.50
232	PHE	N	66.128	43.392	44.016	20.26
234	PHE	CA	67.429	43.035	44.653	17.74
	PHE	C	68.199	42.026	43.725	19.63
235		0	67.571	41.347	42.920	18.88
236	PHE	_	67.148	42.482	46.088	15.74
237	PHE	CB		43.443	47.024	14.99
238	PHE	CG	66.400		47.02 <del>4</del> 47.475	14.29
239	PHE	CD1	67.009	44.602	47.473 47.458	11.07
240	PHE	CD2	65.081	43.181		13.28
241	PHE	CE1	66.294	45.491	48.252	
242	PHE	CE2	64.379	44.048	48.267	12.04
243	PHE	CZ	64.998	45.216	48.653	14.86
244	PRO	N <sub>1</sub>	69.545	41.908	43.722	19.03
245	PRO	CA	70.154	40.759	43.057	17.74
246	PRO	C	69.576	39.346	43.319	17.59
247	PRO	0	69.475	38.926	44.441	18.11
248	PRO	CB	71.601	40.765	43.541	17.29
249	PRO	CG	71.547	41.671	44.730	17.67
250	PRO	CD	70.462	42.691	44.470	17.97
251	CYS	N	69.285	38.589	42.224	16.91
252	CYS	CA	69.204	37.088	42.281	16.83
253	CYS	. C	70.495	36.319	41.751	17.10
254	CYS	0	70.417	35.519	40.790	15.98
255	CYS	CB	68.007	36.580	41.502	17.42
256	CYS	SG	66.580	37.626	41.691	21.76
257	ARG	N	71.667	36.666	42.348	17.15
258	ARG	CA	72.970	36.242	41.791	19.32
259	ARG	С	73.119	34.699	41.788	19.49
260	ARG	0	73.496	34.047	40.814	19.32
261	ÁRG	CB	74.079	36.850	42.622	24.18
262	ARG	CG	74.980	37.833	41.912	34.75
263	ARG	CD	75.311	39.152	42.663	42.80
264	ARG	NE	75.471	39.097	44.113	51.13
265	ARG	CZ	74.828	39.887	45.009	51.47
266	ARG	NH1	74.701	41.151	44.646	48.86



267	ARG	NH2	74.350	39.441	46.136	48.27
268	VAL	N	72.830	34.041	42.924	17.97
269	VAL	CA	72.998	32.576	42.948	16.80
270	VAL	C	71.992	31.806	41.998	15.05
271	VAL	0	72.306	30.779	41.377	15.45
272	VAL	СВ	73.273	32.034	44.312	18.01
			72.573	30.756	44.520	15.99
273	VAL	CG1		32.942	45.388	17.07
274	VAL	CG2	72.876	32.430	41.831	14.53
275	ALA	N	70.819			
276	ALA	CA	69.795	31.819	41.038	14.43
277	ALA	C .	70.213	31.697	39.659	15.46
278	ALA	0	69.747	30.800	38.995	16.10
279	ALA	СВ	68.535	32.680	41.126	12.61
280	LYS	N	71.076	32.655	39.323	17.52
281	LYS	CA	71.665	32.762	37.982	17.70
282	LYS	C	72.998	32.019	37.847	19.60
283	LYS	0	73.580	32.004	36.776	22.60
284	LYS	CB	71.858	34.261	37.582	19.24
285	LYS	CG	70.573	35.200	37.622	19.97
286	LYS	CD	69.498	34.765	36.639	21.69
287	LYS	CE	68.278	35.680	36.497	24.74
288	LYS	ΝZ	67.268	34.860	35.757	26.46
289	LEU	N ·	73.581	31.431	38.936	20.39
290	LEU	CA	74.928	30.843	38.665	19.13
291	LEU	С	74.792	29.677	37.700	21.96
292	LEU	0	73.908	28.875	37.880	20.82
293	LEU	СВ	75.585	30.242	39.923	18.89
294	LEU	CG	75.924	31.268	40.972	17.87
295	LEU	CD1	76.848	32.313	40.400	19.14
296	LEU	CD2	76.506	30.672	42.266	17.39
297	PRO	N	75.721	29.419	36.742	23.61
298	PRO	CA	75.299	28.411	35.767	24.44
299	PRO	C	75.014	27.017	36.391	22.23
300	PRO	Ö	74.374	26.171	35.790	23.82
301	PRO	СВ	76.375	28.491	34.687	26.16
302	PRO	CG	76.896	29.926	34.822	27.75
303		CD	76.869	30.196	36.324	26.72
	PRO				37.638	22.30
304	LYS	N	75.512	26.800	38.233	22.26
305	LYS	CA	75.166	25.529		22.20
306	LYS	С	73.654	25.317	38.489	
307	LYS	0	73.187	24.168	38.569	25.95
308	LYS	CB	76.067	25.121	39.358	23.13
309	LYS	ĊG	75.991	25.832	40.660	24.19
310	LYS	CD	76.895	25.038	41.664	27.45
311	LYS	CE	77.506	25.823	42.852	31.65
312	LYS	NZ	77.983	24.871	43.905	37.10

040	A CAL	N.I	72.000	26 422	38.604	20.49
313	ASN	N	72.908	26.422		
314	ASN	CA	71.547	26.288	39.034	18.95
315	ASN	С	70.578	26.384	37.886	19.83
316	ASN	0	69.364	26.402	38.110	16.61
317	ASN	CB	71.326	27.301	40.083	15.32
318	ASN	CG	72.319	27.082	41.210	18.08
319	ASN	OD1	72.774	25.998	41.593	16.89
320	ASN	ND2	72.630	28.223	41.762	17.04
321	LYS	N	71.115	26.261	36.653	19.50
322	LYS	CA	70.198	26.533	35.517	20.65
323	LYS	C	68.970	25.582	35.499	17.99
324	LYS	0	67.827	25.897	35.252	17.79
325	LYS	СВ	71.102	26.415	34.280	24.77
				27.324	33.113	35.23
326	LYS	CG	70.705			43.17
327	LYS	CD	71.780	27.296	31.993	
328	LYS	CE	71.506	28.319	30.877	48.79
329	LYS	NZ	72.095	27.908	29.585	51.89
330	ASN	N	69.249	24.336	35.828	16.87
331	ASN	CA	68,139	23.356	35.844	17.37
332	ASN	С	67.428	23.162	37.209	14.52
333	ASN	0	67.042	22.047	37.553	14.87
334	ASN	CB	68.758	22.008	35.522	17.56
335	ASN	CG	69.723	21.326	36.476	21.16
336	ASN	OD1	70.099	21.716	37.568	23.28
337	ASN	ND2	70.132	20.210	35.940	22.37
338	ARG	N	67.435	24.272	37.960	13.45
339	ARG	CA	66.851	24.363	39.265	14.08
340	ARG	C	65.834	25.455	39.226	11.72
		0	64.908	25.455	39.985	11.90
341	ARG		67.927	24.635	40.349	14.72
342	ARG	CB			40.637	15.03
343	ARG	CG	68.808	23.399		13.03
344	ARG.	CD	69.531	23.566	41.958	
345	ARG	NE	70.329	22.382	42.127	14.96
346	ARG	CZ	70.786	22.027	43.336	15.87
347	ARG	NH1	70.387	22.635	44.429	14.76
348	ARG	NH2	71.629	21.021	43.409	16.81
349	ASN	N	65.990	26.376	38.305	12.70
350	ASN	CA	64.823	27.286	38.125	12.44
351	ASN	С	63.866	26.703	37.067	12.58
352	ASN	0	64.316	26.017	36.196	11.79
353	ASN	СВ	65.333	28.584	37.553	15.34
354	ASN	CG	66.254	29.036	38.579	15.14
355	ASN	OD1	65.853	29.008	39.721	13.40
356	ASN	ND2	67.478	29.375	38.192	15.01
357	ARG	N	62.602	26.959	37.101	11.31
					36.097	10.21
358	ARG	CA	61.750	26.422	30.087	10.21

359	ARG	С	61.344	27.550	35.032	11.77
360	ARG	0	60.942	27.244	33.909	10.43
361	ARG	CB	60.569	25.882	36.926	10.53
362	ARG	CG	59.435	25.507	35.982	9.14
363	ARG	CD	58.083	25.567	36.545	10.01
364	ARG	NE	57.805	24.437	37.372	10.47
365	ARG	CZ	57.604	23.255	36.784	10.78
366	ARG	NH1	57.345	22.981	35.539	11.53
367	ARG	NH2	57.847	22.305	37.582	10.00
368	TYR	N	61.531	28.839	35.419	12.06
369	TYR	CA	61.406	29.958		10.98
370	TYR	С	62.644	30.880	34.512	- 13.59
371	TYR	Ō	63.313	31.085		15.53
372	TYR	CB	60.188	30.734		
373	TYR	CG	58.902	29.947		11.54
374	TYR	CD1	58.528	29.222		11.75
375	TYR	CD2	58.104	29.917		10.70
376	TYR	CE1	57.373	28.429		12.58
377	TYR	CE2	56.968	29.113	35.885	11.42
378	TYR	CZ	56.577	28.348	34.738	12.61
379	TYR	ОН	55.482	27.467	34.686	13.20
380	ARG	N	63.021	31.367	33.315	13.40
381	ARG	CA	64.268	32.148	33.315	16.28
382	ARG	С	63.947	33.436	34.160	14.00
383	ARG	0	64.804	34.093	34.720	16.44
384	ARG	CB	64.600	32.319	31.810	17.48
385	ÄRG	CG	65.513	33.475	31.434	23.38
386	ARG	CD	65.168	34.185	30.101	28.56
387	ARG	NE	63.860	34.904		
388	ARG	CZ	63.004	34.975	29.117	
389	ARG	NH1	63.482	34.836		
390	ARG	NH2	61.715	35.089		
391	ASP	N	62.671	33.824		
392	ASP .	CA	62.305	35.149		
393	ASP	C	61.714	35.183		
394	ASP	0	61.194	36.232		
395	ASP	CB	61.233	35.719		
396	ASP	CG	61.768	36.276		
397	ASP	OD1	62.985	36.503		
398	ASP	OD2	60.953	36.507		
399	VAL	N	61.838	34.020		
400	VAL	CA	61.386	33.871	38.121	11.29
401	VAL	C	62.363	33.150		
402	VAL	0	62.300	31.998		
403	VAL	CB	59.862	33.736		
404	VAL	CG1	59.564	33.071	39.708	15.44

405 406	VAL SER	CG2 N	59.003 63.292	33.367 34.043	37.026 39.508	14.33 10.16
407	SER	CA	64.351	33.658	40.430	11.16
408	SER	C	64.044	34.235	41.838	9.64
409	SER	0	63.282	35.201	41.980 39.868	11.51 10.81
410	SER	CB	65.674	34.207	38.393	12.18
411	SER	OG	65.795	34.206 33.603	42.890	8.67
412	PRO	N CA	64.652 64.679	34.069	44.300	10.79
413	PRO PRO	CA	65.739	35.174	44.607	13.35
414 415	PRO	0	66.863	35.054	44.167	14.81
416	PRO	CB	65.070	32.742	45.027	10.00
417	PRO	CG	65.954	31.994	44.070	10.77
418	PRO	CD	65.350	32.343	42.713	9.16
419	PHE	N.	65.333	36.312	45.288	13.59
420	PHE	CA	66.396	37.309	45.592	12.71
421	PHE	C	67.463	36.637	46.469	12.31
422	PHE	Ö	67.099	35.793	47.313	12.68
423	PHE	СВ	65.827	38.538	46.324	9.81
424	PHE	CG	64.721	4 39.284	45.705	10.06
425	PHE	CD1	64.796	39.648	44.360	9.58
426	PHE	CD2	63.667	39.689	46.481	9.41
427	PHE	CE1	63.839	40.478	43.828	10.26
428	PHE	CE2	62.712	40.531	45.935	10.02
429	PHE	CZ	62.810	40.954	44.617	8.64
430	ASP	N ·	68.731	37.039	46.301	11.62
431	ASP	CA	69.699	36.442	47.229	12.07
432	ASP	С	69.336	36.779	48.740	12.27
433	ASP	0	69.471	35.939	49.614	15.21
434	ASP	CB	71.090	36.960	46.918	13.40
435	ASP	CĢ	71.585	36.819	45.512	14.25
436	ASP	OD1	71.510	35.755	45.001	15.72
437	ASP	OD2	72.114	37.776	44.954	12.79
438		N	68.891	37.987	49.110	12.16
439	HIS	CA	68.936	38.332	50.537	12.71
440	HIS	C	68.037	37.415	51.373	13.92
441	HIS	0	68.357	37,023	52.468	15.61
442	HIS	CB	68.619	39.807	50.639	12.05
443	HIS	CG	67.189	40.229	50.532	10.80
444	HIS	ND1	66.218	40.158	51.532	11.89
445	HIS	CD2	66.595	40.810	49.442	10.89
446	HIS	CE1	65.078	40.670	51.055	9.38
447	HIS	NE2	65.292	41.065	49.801	11.30 13.43
448	SER	N 	66.911	37.032	50.727 51.219	12.13
449	SER	CA	65.732	36.268 34.786	51.219	12.13
450	SER	С	65.616	34.700	JU.142	12.70

451	SER	0	64.852	33.988		51.280		13.37
452	SER	СВ	64.459	36.969		50.655		10.16
453	SER	OG	64.073	36.712		49.237		10.34
454	ARG	N	66.289	34.396		49.662	•	11.78
455	ARG	CA	66.110	33.011		49.184	,	13.97
456	ARG	C	66.308	32.030		50.330		13.85
457	ARG	0	67.052	32.388		51.228		11.86
		CB	67.182	32.758		48.072		12.96
458	ARG			32.736		48.534		13.81
459	ARG	CG	68.663			47.602		13.20
460	ARG	CD	69.587	31.920				14.78
461	ARG	NE	70.951	31.816		48.184		
462	ARG	CZ	71.552	30.688		48.524	•	15.00
463	ARG	NH1	70.936	29.557		48.478		11.82
464	ARG	NH2	72.784	30.639		48.884		17.64
465	ILE	Ν	65.654	30.828		50.300		14.21
466	ILE	CA	65.948	29.645		51.217		12.58
467	ILE	С	67.155	28.748	•	50.720		14.61
468	ILE	0	67.216	28.241		49.587		15.26
469	ILE	CB	64.722	28.774		51.313		13.58
470	ILE	CG1	63.510	29.500		51.879		13.29
471	ILE	CG2	64.977	27.447		52.006		13.64
472	ILE	CD1	63.571	29.924		53.338		13.30
473	LYS	N	68.150	28.621		51.612		15.82
474	LYS	CA	69.276	27.764		51.318		15.98
475	LYS	С	69.096	26.360		51.954		17.60
476	LYS	0	68.984	26.265		53.135		19.95
477	LYS	СВ	70.448	28.373		52.073		17.28
478	LYS	CG	70.639	29.851		51.912		17.24
479	LYS	CD	72.028	30.137		52.423		22.81
480	LYS	CE	72.374	31.627		52.475		26.05
481	LYS	NZ	73.787	31.719		52.904		31.86
482	LEU	N	69.186	25.284		51.179		17.25
483	LEU	CA	69.191	23.986		51.803		18.29
484	LEU	C	70.420	23.813		52.707		20.34
	LEU	0	71.463	24.451		52.542		19.70
485				23.009		50.654		18.28
486	LEU	CB	69.326			49.875		18.66
487	LEU	CG	68.077	22.606			*	
488	LEU	CD1	68.503	22.354		48.425		17.13
489	LEU	CD2	66.905	23.584		50.003		14.98
490	HIS	N	70.327	22.853		53.607		22.86
491	HIS	CA	71.473	22.568		54.469		25.86
492	HIS	С	72.376	21.544		53.818		30.06
493	HIS	0	72.823	20.636		54.478		31.82
494	HIS	CB	71.011	22.101		55.889		26.12
495 -	-HIS	CG	-70-169	 -23:121		56.652		25.12
496	HIS	ND1	69.669	22.796		57.857		25.76

497	HIS	CD2	69.771	24.443	56.370	24.91
498	HIS	CE1	68.999	23.876	58.293	25.52
499	HIS	NE2	69.054	24.883	57.439	26.33
500	GLN	N	72.654	21.694	52.528	32.94
501	GLN	CA	73.476	20.669	51.845	36.50
502	GLN	С	74.845	21.250	51.490	37.97
503	GLN	0	74.997	22.462	51.350	37.18
504	GLN	СВ	72.785	19.969	50.653	37.95
505	GLN	CG	72.220	20.893	49.533	40.06
506	GLN	CD	71.786	20.148	48.187	44.06
507	GLN	OE1	72.396	20.225	47.103	46.49
508	GLN	NE2	70.689	19.404	48.371	41.49
509	GLU	N	75.833	20.321	51.371	41.15
510	GLU	CA	77.175	20.785	50.982	41.70
511	GLU	C	77.356	20.946	49.420	39.31
512	GLU	Ö	78.061	21.828	48.944	38.68
513	GLU	СВ	78.182	19.780	51.511	45.57
514	GLU	CG	78.185	19.584	53.033	54.22
515	GLU	CD	78.719	18.188	53.518	60.23
516	GLU	OE1	79.014	17.293	52.706	63.36
517	GLU	OE2	78.813	18.019	54.741	62.48
518	ASP	N	76.708	20.105	48.601	38.53
519	ASP	CA	77.055	20.312	47.192	37.73
520	ASP	C	76.579	21.706	46.692	33.50
521	ASP	Ö	77.309	22.594	46.250	35.46
522	ASP	СВ	76.361	19.144	46.491	43.21
523	ASP	CG	76.711	19.135	45.008	49.96
524	ASP	OD1	77.838	19.533	44.702	53.55
525	ASP	OD2	75.843	18.784	44.184	52.89
526	ASN	N	75.262	21.863	46.853	28.37
527	ASN	CA	74.553	23.027	46.358	21.99
528	ASN	C	73.354	23.441	47.299	20.24
529	ASN	Ö	72.312	22:805	47.353	21.87
530	ASN	СВ	74.099	22.536	44.960	19.04
531	ASN	CG	73.717	23.704	44.039	18.04
532	ASN	OD1	73.456	24.765	44.530	19.25
533	ASN	ND2	73.728	23.588	42.734	15.06
534	ASP	N.	73.466	24.546	48.022	18.00
535	ASP	CA	72.332	25.033	48.836	18.78
536	ASP	C	71.151	25.681	48.030	17.35
537	ASP	0	70.225	26.206	48.623	18.81
538	ASP	СВ	72.886	26.063	49.861	19.48
539	ASP	CG	73.288	27.453	49.333	23.10
540	ASP	OD1	72.887	27.851	48.256	24.35
541	ASP	OD2	73.983	28.207	50.021	29.58
542	TYR	N:	71.212	25.740	46.675	16.55

543	TYR	CA	70.199	26.452	45.895	12.98
544	TYR	С	68.875	25.687	45.783	12.27
545	TYR	Ō	68.835	24.576	45.292	12.81
546	TYR	СВ	70.772	26.662	44.525	10.36
547	TYR	ĊĠ	69.834	27.547	43.751	12.42
548	TYR	CD1	69.670	28.900	44.065	10.55
549	TYR	CD2	69.026	27.003	42.736	11.80
550	TYR	CE1	68.666	29.636	43.411	11.79
551	TYR	CE2	68.002	27.711	42.094	10.71
552	TYR	CZ	67.821	29.072	42.453	9.67
553	TYR	OH	66.838	29.931	41.987	10.99
554	ILE	N	67.843	26.362	46.233	13.06
555	ILE	CA	66.470	26.037	45.852	12.41
556	ILE	C	65.728	27.348	45.344	11.32
	ILE	0	66.023	28.445	45.836	12.19
557 559		CB	65.643	25.321	46.943	12.13
558	ILE		64.216	25.028	46.401	8.76
559	ILE	CG1		26:109	48.238	10.91
560	ILE	CG2	65.631		46.236	9.88
561	ILE	CD1	63.534	23.768	44.373	9.16
562	ASN	N	64.802	27.274		
563	ASN	CA	63.961	28.432	44.113	9.65
564	ASN	C	62.777	28.477	45.134	10.37
565	ASN	0	61.669	28.001	44.902	10.07
566	ASN	СВ	63.409	28.368	42.689	9.08
567	ASN	CG	62.854	29.669	42.094	9.54
568	ASN	OD1	62.064	30.411	42.717	10.96
569	ASN	ND2	63.215	29.836	40.823	9.06
570	ALA	N	63.053	29.245	46.232	9.41
571	ALA	CA	62.074	29.518	47.285	10.05
572	ALA	С	62.490	30.729	48.081	11.58
573	ALA	0	63.684	31.012	48.161	12.92
574	ALA	СВ	62.248	28.369	48.268	7.50
575	SER	N '	61.501	31.398	48.688	10.51
576	SER	CA	61.767	32.622	49.448	11.23
577	SER	C	60.986	32.733	50.725	10.85
578	SER	Ô	59.810	32.408	50.715	12.48
579	SER	CB	61.240	33.808	48.570	7.55
580	SER	OG	61.832	33.943	47.206	10.45
581	LEU	N	61.665	33.227	51.759	11.48
582	LEU.	CA	61.028	33.484	53.026	. 12.53
583	LEU	С	60.459	34.863	52.839	13.15
584	LEU	0	61.162	35.810	52.745	13.65
585	LEU	СВ	62.100	33.317	54.125	13.18
586	LEU	CG	61.714	33.026	55.617	15.58
587	LEU	CD1	60.346	32.465	56.032	15.45
588	LEU	CD2	61.990	34.242	56.446	15.90

589	ILE	N	59.142	34.963	52.710	12.91
590	ILE	CA	58.433	36.244	52.854	13.87
591	ILE	C	58.199	36.410	54.383	16.47
592	ILE	Ö	57.520	35.626	55.029	16.73
593	ILE	СВ	57.078	36.103	52.126	13.55
594	ILE	CG1	57.058	36.090	50.580	12.82
595	ILE	CG2	56.084	37.173	52.640	16.67
596	ILE	CD1	58.184	35.285	49.937	12.75
597	LYS	N	58.754	37.425	54.954	16.38
598	LYS	CA	58.733	37.584	56.393	19.03
599	LYS	C	57.916	38.848	56.792	18.79
600	LYS	Ö	58.361	39.997	56.871	18.39
601	LYS	CB	60.193	37.663	56.758	24.89
602	LYS	CG	60.382	37.380	58.215	36.70
603	LYS	CD	61.837	37.012	58.412	45.35
604	LYS	CE	62.107	36.507	59.815	49.89
605	LYS	NZ	63.542	36.156	59.861	53.29
606	MET	Ν	56.632	38.598	57.069	17.44
607	MET	CA	55.789	39.713	57.508	17.07
608	MET	С	56.013	39.963	59.034	17.04
609	MET	0	55.491	39.236	59.879	16.87
610	MET	CB	54.344	39.363	57.116	15.46
611	MET	C.G	54.239	38.984	55.611	15.47
612	MET	SD	55.075	40.191	54.531	16.71
613	MET	CE	53.670	41.304	54.490	12.89
614	GLU	N	56.792	41.029	59.352	20.27
615	GLU	CA	57.059	41.404	60.754	22.54
616	GLU	С	55.827	42.015	61.552	22.33
617	GLU	0	55.304	41.421	62.490	24.08
618	GLU	CB	58.262	42.322	60.762	26.36
619	GLU	CG	58.891	42.457	62.173	35.25
620	GLU	CD	59.916	43.590	62.166	40.00
621	GLU	OE1	59.477	44.700	61.830	43.15
622	GLU	OE2	61.096	43.345	62.480	43.54
623	GLU	N	55.283	43.165	61.103	23.41 24.81
624	GLU	CA	54.028	43.623	61.715	24.01
625	GLU	C	52.959	42.499	61.936	24.74
626	GLU	0	52.437	42.295 44.704	63.015 60.837	26.99
627	GLU	CB	53.482 52.576	45.679	61.608	35.79
628	GLU	CG CD	51.974	46.718	60.634	45.06
629	GLU	OE1	52.430	46.765	59.511	48.32
630 631	GLU	OE2	51.091	47.475	61.022	49.68
632	ALA	N	52.665	41.750	60.868	22.17
633	ALA	CA	51.673	40.706	60.947	21.64
634	ALA	C	52.213	39.523	61.660	22.46
JJ7		•	JZ.Z 1 J	00.020	31.300	



635	ALA	0	51.470	38.690	62.105	24.05
636	ALA	СВ	51.310	40.191	59.565	20.53
637	GLN	N	53.508	39.446	61.797	23.74
638	GLN	CA	53.951	38.375	62.649	27.80
639	GLN	C	53.494	36.942	62.164	26.92
640	GLN	Ō	53.212	36.073	62.988	27.71
641	GLN	СВ	53.604	38.695	64.142	32.88
642	GLN -	CG	54.557	39.626	64.990	40.36
643	GLN	CD	55.637	38.789	65.755	47.19
644	GLN	OE1	55.413	38.194	66.807	52.84
645	GLN	NE2	56.809	38.690	65.144	46.59
646	ARG	N	53.576	36.814	60.781	23.43
647	ARG	CA	53.512	35.548	59.983	18.91
648	ARG	C	54.532	35.508	58.799	16.84
649	ARG	Ö	54.660	36.485	58.081	17.96
650	ARG	CB	52.108	35.410	59.417	18.58
651	ARG	CG	51.983	33.974	58.874	16.36
652	ARG	CD	50.561	33.599	58.555	17.70
653	ARG	NE	49.917	33.327	59.836	17.29
654	ARG	CZ	48.629	33.215	59.998	15.94
655	ARG	NH1	47.869	33.433	58.973	13.68
656	ARG	NH2	48.184	32.917	61.156	18.40
657	SER	N	55.331	34.440	58.582	13.82
658	SER	CA	56.164	34.277	57.366	12.48
659	SER	C	55.588	33.151	56.471	11.37
660	SER	Ö	54.899	32.287	56.921	13.16
661	SER	CB.	57.601	33.787	57.603	12.86
662	SER	ŌĞ	58.317	33.977	58.878	19.01
663	TYR	N	56.004	33.108	55.215	11.99
664	TYR	CA	55.704	31.895	54.458	10.12
665	TYR	C	56.953	31.593	53.701	10.62
666	TYR	Ō	57.730	32.497	53.459	11.32
667	TYR	СВ	54.616	32.184	53.384	10.39
668	TYR	CG	53.469	33.063	53.849	10.08
669	TYR.	CD1	53.696	34.400	54.011	10.55
670	TYR	CD2	52.208	32.580	54.154	10.53
671	TYR	CE1	52.769	35.233	54.522	12.90
672	TYR	CE2	51.239	33.420	54.642	12.10
673	TYR	CZ	51.530	34.723	54.834	13.13
674	TYR	ОН	50.524	35.465	55.346	13.22
675	ILE	N	57.104	30.369	53.235	10.35
676	ILE	CA	58.077	30.153	52.147	7.97
677.	ILE,	C	57.178	30.151	50.851	9.78
678	ILE	0	56.263	29.348	50.833	10.25
679-	TLE	CB -	58.953	28.883	52.458	9.05
680	ILE	CG1	59.740	29.088	53.740	10.20
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681 682 683 684 685 686	ILE ILE LEU LEU LEU LEU	CG2 CD1 N CA C	59.957 60.474 57.403 56.855 57.905 59.110	28.496 27.819 31.084 30.922 30.236 30.576	51.397 54.246 49.833 48.465 47.526 47.552	8.37 8.33 8.73 8.66 11.10 12.49
687 688	LEU LEU	CB CG	56.561 55.257	32.244 32.826	47.860 48.326	9.65 10.58
689	LEU	CD1	54.913	34.147	47.565	11.19
690	LEU	CD2	55.270	32.940	49.838	12.47
691	THR	N	57.402	29.203	46.761	10.83
692	THR	CA	58.299	28.363	45.895	8.06
693	THR	С	57.668	28.097	44.482	9.10
694	THR	0	56.489	28.380	44.264	8.53
695	THR	CB	58.859	27.100	46.647	8.87
696	THR	OG1	59.976	26.366	45.972	11.27
697	THR	CG2	57.803	26.211	47.344	8.63
698	GLN	Ν	58.506	27.610	43.519	7.55
699	GLN	CA	57.990	27.172	42.176	7.51
700	GLN	. C	57.437	25.750	42.219	8.15
701	GLN	0	57.723	24.997	43.143	9.26
702	GLN	CB	59.060	27.244	41.087	7.72
703	GLN	CG	60.130	26.188	41.317	7.48 7.54
704	GLN -	CD OE1	61.257	26.336 25.398	40.384 40.166	13.65
705 706	GLN GLN	NE2	61.977 61.511	27.505	39.888	6.16
700	GLN	NEZ	`56.603	25.377	41.221	8.70
707	GLY	CA	56.238	23.982	41.272	8.95
709	GLY	C	57.520	23.130	41.108	10.89
710	GLY	Ö	58.254	23.333	40.156	12.56
711	PRO	Ň	57.762	22.179	42.005	11.90
712	PRO	CA	58.999	21.445	41.898	10.45
713	PRO	С	59.179	20.901	40.444	12.59
714	PRO	0	58.192	20.592	39.776	12.30
715	PRO	CB	58.889	20.400	42.984	10.51
716	PRO	CG	57.757	20.889	43.873	11.89
717	PRO	CD	56.917	21.895·	43.147	9.07
718	LEU	N	60.423	20.963	39.954	13.19
719	LEU	CA	60.928	20.422	38.711	14.02
720	LEU	С	61.288	18.972	38.994	13.93
721	LEU	0	61.401	18.617	40.157	12.95
722	LEU	CB	62.178	21.303 -	38.472	13.39
723	LEU.		62.181	22.123	37.185	14.69
724	LEU	CD1	62.862	23.432	37.390 36.589	13.28 12.21
725	PRO	CD2 N	60.808	22.403 18.060	38.006	16.81
<del>7</del> 26	FKU	IN	01.482	10.000	30.000	10.01

707	DDO	CA	61.666	16.641	38.394	17.34
727	PRO				39.104	16.85
728	PRO	С	62.962	16.281	39.688	18.24
729	PRO	0	63.122	15.247		
730	PRO	CB	61.517	15.864	37.116	17.12
731	PRO	CG	60.922	16.848	36.100	19.47
732	PRO	CD	61.300 <sub>,</sub>	18.261	36.556	16.45
733	ASN	N	63.900	17.198	39.011	15.30
734	ASN	CA	65.204	16.949	39.638	14.86
735	ASN	C	65.487	17.673	40.952	15.36
736	ASN	0	66.603	17.703	41.410	15.84
737	ASN	CB	66.197	17.452	38.620	14.48
738	ASN	CG	65.975	18.909	38.203	16.03
739	ASN	OD1	65.058	19.216	37.443	1,9.68
740	ASN	ND2	66.878	19.774	38.721	15.67
741	THR	N	64.484	18.374	41.425	15.02
742	THR	CA	64.442	19.135	42.712	12.29
743	THR	C	63.211	18.662	43.532	11.96
744	THR	0	62.712	19.350	44.385	11.69
745	THR	СВ	64.049	20.608	42.328	10.87
745	THR	OG1	62.724	20.721	41.769	10.92
		CG2	64.891	21.218	41.196	10.32
747	THR				43.321	14.04
748	CYS	N	62.634	17.505	44.179	14.17
749	CYS	CA	61.483	17.193		
750	CYS	C	62.088	16.768	45.539	12.97
751	CYS	0	61.464	16.855	46.615	13.90
752	CYS	CB	60.644	16.031	43.646	13.16
753	CYS	SG	59.565	16.488	42.264	15.24
754	GLY	N	63.405	16.394	45.404	13.25
755	GLY	CA	64.191	15.974	46.563	14.11
756	GLY	С	64.788	17.158	47.402	14.77
757	GLY	0	64.846	17.059	48.598	15.71
758	HIS	Ν	65.185	18.303	46.789	15.04
759	HIS	CA	65.415	19.657	47.379	13.49
760	HIS	С	64.101	20.265	47.972	14.38
761	HIS	0	64.083	20.721	49.105	15.19
762	HIS	CB	65.898	20.682	46.344	12.89
763	HIS	CG	67.036	20.181	45.457	14.63
764	HIS	ND1	67.075	20.421	44.117	13.48
765	HIS	CD2	68.117	19.333	45.753	16.55
766	HIS	CE1	68.090	19.729	43.613	14.33
767	HIS	NE2	68.744	19.084	44.580	15.87
768	PHE	N	62.967	20.201	47.261	12.72
769	PHE	CA.	61.732	20.613	47.958	12.68
770	PHE	C	61.548	19.898	49.353	13.22
771	PHE	0	61.483	20.497	50.423	12.12
772	PHE	СВ	60.586	20.347	46.968	12.82
112		<u> </u>	30.000	_0.0 17	.0.000	

773	PHE	CG	59.235	20.806	47.480	9,60
774	PHE	CD1	58.367	19.945	48.151	8.18
775	PHE	CD2	58.819	22.104	47.261	8.54
776	PHE	CE1	57.135	20.383	48.629	11.79
777	PHE	CE2	57.551	22.497	47.662	11.03
778	PHE	CZ	56.712	21.647	48.354	10.70
779	TRP	N	61.531	18.566	49.377	12.17
780	TRP	CA	61.277	17.960	50.667	12.15
781	TRP	C	62.426	18.110	51.675	12.77
782	TRP	Ö	62.267	17.797	52.824	12.89
783	TRP	СВ	60.914	16.499	50.457	13.28
784	TRP	CG	59.487	16.421	49.968	13.46
785	TRP	CD1	59.176	16.003	48.715	13.99
786	TRP	CD2	58.238	16.857	50.571	12.69
787	TRP	NE1	57.858	16.037	48.514	13.76
	TRP	CE2	57.224	16.674	49.611	12.21
788		_		17.354	51.835	11.54
789	TRP	CE3	57.892		49.845	11.49
790	TRP	CZ2	55.928	17.024	· -	12.46
791	TRP	CZ3	56.569	17.707	52.119	11.61
792	TRP	CH2	55.598	17.545	51.099	
793	GLU	N	63.588	18.588	51.263	13.48
794	GLU	CA	64.731	18.763	52.162	12.78
795	GLU	C	64.458	20.064	52.881	13.54
796	GLU	0	64.488	20.065	54.123	13.56
797	GLU	CB	66.084	18.779	51.401	11.88
798	GLU	CG	67.269	19.356	52.235	15.11
799	GLU	CD	68.554	19.518	51.499	17.43
800	GLU	OE1	68.577	19.290	50.284	19.12
801	GLU	OE2	69.547	19.872	52.101	19.81
802	MET	Ν.,	63.973	21.049	51.994	11.71
803	MET	CA	63.546	22.379	52.486	12.23
804	MET	С	62.389	22.312	53.511	11.68
805	MET	0	62.291	22.943	54.555	11.24
806	MET	CB	63.108	23.239	51.328	9.82
807	MET	CG	62.214	24.420	51.724	11.63
808	MET	SD	61.999	25.602	50.392	15.04
809	MET	CE	60.555	24.934	49.564	11.38
810	VAL	N	61.422	21.461	53.168	12.17
811	VAL	CA	60.354	21.227	54.164	10.63
812	VAL	C	60.941	20.624	55.438	13.77
813	VAL	Ō	60.589	21.011	56.523	14.26
814	VAL	СВ	59.220	20.388	53.522	10.58
815	VAL	CG1		21.226	52.398	8.22
816	VAL	CG2	58.132	19.960	54.487	10.48
817	TRP	N N	61.857	19.696	55.332	13.52
818	TRP	CA	62.368	19.071	56.546	14.59
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819	TRP	С	63.063		20.108		57.515		15.04 16.22
820	TRP	0	62.665		20.298		58.672		12.97
821	TRP	CB	63.263		17.908		56.111 57.352		17.46
822	TRP	CG	63.663		17.210		58.070		21.61
823	TRP	CD1	64.811		17.506	*			19.79
824	TRP	CD2	62.942		16.235		58.122		
825	TRP	NE1	64.831		16.777		59.222		23.85
826	TRP	CE2	63.721		15.974		59.290		22.70
827	TRP	CE3	61.749	•	15.593		57.954		21.09
828	TRP	CZ2	63.300		15.074		60.220		21.76
829	TRP	CZ3	61.323		14.679		58.916		21.15
830	TRP	CH2	62.094		14.409		60.031		21.23
831	GLU	N	64.079		20.742		56.935		16.06
832	GLU	CA	65.024		21.609		57.615		15.05
833	GLU	C	64.396		22.880		58.008		16.50
834	GLU	0	64.758		23.461		59.019		18.37
835	GLU	CB	66.099		21.948		56.599		13.43
836	GLU	CG	66.738		20.660		56.125		13.77
837	GLU	CD	67.820		20,935		55.146		14.43
838	GLU	OE1	67.899		22.007		54.586		15.87
839	GLU	OE2	68.632		20.038		54.975		15.61
840	GLN	N	63.390		23.283		57.230	•	14.99
841	GLN	CA	62.612		24.477		57.635		14.10
842	GLN	C	61.510		24.226		58.676		14.79
843	GLN	O	60.879		25.146		59.160		14.03
844	GLN	CB	62.048		25.170		56.347		13.84
845	GLN	CG	63.212		25.595		55.375		14.77
846	GLN	CD	64.194		26.544		56.098		20.12
847	GLN	OE1	63.694		27.506		56.665		21.85
848	GLN	NE2	65.471		26.331		56.071		18.11
849	LYS	N	61.251		22.954		59.012	*	14.24
850	LYS	CA	60.227		22.483		59.980		15.01
851	LYS	С	58.797		22.890		59.696		12.96
852	LYS	0	57.994		22.992		60.604		13.31
853	LYS	CB	60.584		22.912	•	61.402		19.21
854	LYS	CG	62.055		22.609		61.711		21.85
855	LYS	CD	62.235		22.379	,	63.198		29.47
856	LYS	CE	63.661		22.216		63.640		31.82
857	LYS	NZ	64.341		23.105		62.727		37.86
858	SER	N	58.495		23.079		58.406		14.72
859	SER	CA	57.092		23.282		58.019		14.22
860	SER	C	56.246		22.042		58.300		15.12
861 .	SER	0	56.655		20.897		58.189		14.48
862	SER	CB	57.124		23.649		56.489		12.78
863	SER	OG	58.109		24.701		56.039		13.54
864	ARG	N	55.046	•	22.316		58.638		13.87

865	ARG	CA	53.889	21.469	58.787	14.62
866	ARG	C	52.900	21.380	57.558	14.77
867	ARG	Ö	52.279	20.349	57.295	13.72
868	ARG	СВ	53.162	22.084	59.989	14.39
869	ARG	ČG	52.448	21.021	60.790	23.02
870	ARG	CD	51.000	20.993	60.443	27.49
	ARG	NE	50.286	20.242	61.462	30.20
871		CZ	50.380	18.916	61.522	29.90
872	ARG	NH1	•	18.211	60.950	31.36
873	ARG		51.307		62.133	30.49
874	ARG	NH2	49.427	18.304	56.827	15.17
875	GLY	N	52.739	22.450		13.17
876	GLY	CA	51.776	22.626	55.799	
877	GLY	С	52.462	22.850	54.485	12.93
878	GLY	0	53.449	23.538	54.399	12.18
879	VAL	N	51.894	22.272	53.426	13.38
880	VAL	CA	52.274	22.713	52.082	10.87
881	VAL	C	50.983	23.124	51.461	9.66
882	VAL	0	50.002	22.387	51.541	10.04
883	VAL ?	CB	52.870	21.528	51.228	10.73
884	VAL	CG1	54.155	20.840	51.773	11.72
885	VAL	CG2	53.136	21.897	49.737	10.72
886	VAL	N	51.007	24.297	50.825	8.51
887	VAL	CA	49.805	24.732	50.087	7.91
888	VAL	C	50.113	24.799	48.608	8.39
889	VAL	0	50.967	25.546	48.186	9.71
890	VAL	СВ	49.346	26.112	50.615	6.14
891	VAL	CG1	48.848	26.046	52.098	7.70
892	VAL	CG2	48.214	26.748	49.800	6.32
893	MET	.N	49.389	24.005	47.841	8.82
894	MET	CA	49.534	23.899	46.386	8.53
895	MET	С	48.365	24.505	45.638	9.29
896	MET	Ō	47.289	23.983	45.711	9.66
897	MET	CB :	49.534	22.412	46.115	8.63
898	MET	CG	49.584	22.144	44.622	9.02
899	MET	SD	50.458	20.629	44.313	13.59
900	MET	CE	50.211	20.435	42.521	7.30
901	LEU	N.	48.591	25.598	44.954	8.05
902	LEU	CA	47.428	26.275	44.340	6.79
903	LEU	C	47.172	26.020	42.785	8.29
904	LEU	0	46.566	26.877	42.182	8.17
	LEU		47.688	27.800	44.449	8.82
905		CB		28.331	45.842	9.33
906	LEU	CG	47.991			6.68
907	LEU	CD1	46.842		46.813	8.69
908	LEU	CD2	48.214	29.832	45.771	8.85
909	ASN	- N	47.920	25.017	42.200	
910	ASN	CA	48.020	24.702	40.735	10.85

911	ASN	С	47.769	23.175	40.514	10.96
912	ASN	0	48.012	22.421	41.451	11.52
913	ASN	CB	49.393	25.060	40.117	10.56
914	ASN	CG	50.568	24.084	40.487	10.18
915	ASN	OD1	51.127	23.276	39.749	13.30
916	ASN	ND2	51.129	24.353	41.654	6.84
917		ND2 N	47.424	22.750	39.274	12.81
	ARG	CA	47.424	21.308	38.970	12.84
918	ARG		48.776	20.954	38.393	13.06
919	ARG	С		21.835	37.869	13.72
920	ARG	O	49.452		38.085	15.72
921	ARG	CB	46.231	20.948		23.73
922	ARG	CG	44.960	21.740	38.453	
923	ARG	CD	43.614	21.096	38.015	34.40
924	ARG	NE	43.483	20.691	36.595	44.56
925	ARG	CZ	43.334	21.619	35.619	51.42
926	ARG	NH1	43.467	22.892	35.787	54.99
927	ARG	NH2	42.994	21.256	34.403	53.99
928	VAL	N	49.195	19.695	38.452	13.43
929	VAL	CA	50.346	19.256	37.704	16.12
930	VAL	С	50.278	19.565	36.147	16.48
931	VAL	0	51.114	20.226	35.584	14.56
932	VAL	CB	50.559	17.783	38.070	14.79
933	VAL	CG1	50.930	17.661	39.553	14.80
934	VAL	CG2	51.639	17.182°	37.160	16.58
935	MET	N	49.229	19.073	35.453	17.62
936	MET	CA	48.799	19.616	34.173	19.41
937	MET	С	47.577	20.606	34.287	17.22
938	MET	0	46.471	20.315	34.733	16.41
939	MET	CB	48.548	18.474	33.180	22.53
940	MET	CG	48.920	18.893	31.711	30.44
941	MET	SD	48.488	17.626	30.498	36.43
942	MET	CE	46.891	18.345	30.183	32.23
943	GLU	N	47.925	21.768	33.762	17.49
944	GLU	CA	47.082	22.928	33.568	18.64
945	GLU	C	47.552	23.550	32.237	17.88
946	GLU	Ō	48.753	23.611	31.959	18.32
947	GLU	СВ	47.409	24.092	34.512	16.54
948	GLU	CG	47.652	23.731	35.929	16.82
949	GLU	CD	47.619	24.979	36.772	15.94
950	GLU	OE1	48.517	25.821	36.630	15.52
951	GLU	OE2	46.727	25.093	37.611	15.98
952	LYS	N	46.605	23.992	31.444	19.17
953	LYS	CA	46.693	24.610	30.148	20.14
954	LYS	C	47.513	23.739	29.202	18.74
	LIS LYS	. O	48.351	24.244	28.449	18.86
	LYS	СВ	47.304	25.984	30.368	21.50
956	LIS	CD	47.304	20.304	Ç0.000	۲.00

957	LYS	CG	46.267	26.889	31.035	24.11
958	LYS	CD	46.770	28.322	31.089	28.82
959	LYS	CE	45.657	29.354	31.107	30.30
			44.532	28.741	31.820	32.58
960	LYS	NZ		22.392	29.319	18.79
961	GLY	N	47.273		28.344	17.83
962	GLY	CA	47.911	21.493		19.02
963	GLY	C	49.363	21.124	28.602	18.80
964	GLY	0	49.889	20.163	28.022	
965	SER	N	49.994	21.951	29.434	16.08
966	SER	CA	51.417	21.727	29.690	17.76
967	SER	С	51.563	21.319	31.153	16.07
968	SER	0	50.624	21.508	31.921	15.80
969	SER	CB	52.195	22.984	29.281	21.32
970	SER	OG	51.454	23.745	28.215	29.95
971 -	LEU	N	52.706	20.665	31.433	16.19
972	LEU	CA	53.044	20.095	32.721	14.22
973	LEU	С	53.814	21.162	33.523	14.90
974	LEU	0	54.937	21.513	33.212	16.30
975	LEU	CB	53.928	18.847	32.505	13.16
976	LEU	CG	53.098	17.636	32.063	12.66
977	LEU	CD1	52.255	17.105	33.249	15.43
978	LEU	CD2	53.993	16.510	31.569	13.88
979	LYS	N	53.070	21.738	34.484	14.30
980	LYS	CA	53.355	22.962	35.234	11.19
981	LYS	С	54.103	22.727	36.555	10.51
982	LYS	0	54.480	23.691	37.202	11.24
983	LYS	СВ	51.969	23.563	35.492	11.90
984	LYS	CG	51.377	24.141	34.173	15.22
985	LYS	CD	52.251	25.323	33.894	16.86
986	LYS	CE	51.945	26.341	32.815	20.26
987	LYS	NZ	52.785	27.477	33.270	18.45
988	CYS	N.	54.265	21.458	36.919	10.35
989	CYS	CA	54.800	21.085	38.199	10.12
990	CYS	C	55.093	19.570	38.124	13.34
991	CYS	Ö	54.346	18.872	37.449	13.61
992	CYS	CB	53.752	21.438	39.298	13.04
993	CYS	SG	54.171	21.430	41.056	11.06
	ALA	N	56.092	19.102	38.912	11.84
994				17.664	39.115	11.47
995	ALA	CA	56.384			13.20
996	ALA	С	55.350	16.986	39.954	10.87
997	ALA	0	54.725	17.565	40.853	
998	ALA	CB	57.698	17.551	39.874	9.90
999	GLN	N	55.227	15.685		12.25
1000	GLN	CA	54.380	14.965	40.731	13.18
1001	GLN		55.285	14.601	41.907	13.71
1002	GLN	0	55.974	13.592	41.922	14.92

1003	GLN	СВ	53.719	13.735	40.112	13.64
1004	GLN	CG	52.758	12.973	41.060	14.19
1005	GLN	CD	51.427	13.725	41.216	14.78
1006	GLN	OE1	50.905	13.910	42.309	20.52
1007	GLN	NE2	50.901	14.217	40.086	13.59
1008	TYR	N	55.306	15.502	42.891	12.19
1009	TYR	CA	56.373	15.447	43.967	11.14
1010	TYR	C	55.940	14.903	45.328	10.72
1011	TYR	0	56.779	14.953	46.207	12.61
1012	TYR	CB	56.928	16.825	44.297	10.70
1013	TYR	CG	55.892	17.745	44.873	12.19
1014	TYR	CD1	55.710	17.845	46.233	10.21
1015	TYR	CD2	55.141	18.558	44.050	11.16
1016	TYR	CE1	54.801	18.729	46.793	12.37
1017	TYR	CE2	54.213	19.435	44.555	10.83
1018	TYR	CZ	54.033	19.497	45.943	11.09
1019	TYR	ОН	53.050	20.302	46.488	11.70
1020	TRP	N	54.662	14.460	45.484	11.94
1021	TRP	CA	54.108	13.696	46.608	12.84
1022	TRP	С	53.453	12.393	46.058	14.59
1023	TRP	0	53.073	12.404	44.901	14.57
1024	TRP	CB	53.090	14.582	47.358	12.85
1025	TRP	CG	51.796	14.668	46.559	13.65
1026	TRP	CD1	50.660	13.882	46.775	14.58
1027	TRP	CD2	51.462	15.520	45.460	12.43
1028	TRP	NE1	49.663	14.196	45.908	14.02
1029	TRP	CE2	50.123	15.214	45.098	12.63
1030	TRP	CE3	52.186	16.429	44.734	13.32
1031	TRP	CZ2	49.544	15.874	44.043	12.31
1032	TRP	CZ3	51.623	17.068	43.649	11.49
1033	TRP	CH2	50.298	16.797	43.312	14.96
1034	PRO	N	53.291	11.284	46.898	16.00
1035	PRO	CA	52.633	10.039	46.450	15.76
1036	PRO	С	51.096	10.056	46.250	16.40
1037	PRO	0	50.249	10.513	47.021	18.51
1038	PRO	CB	53.112	8.965 47.40		
1039	PRO	CG	53.352	9.784 48.63		
1040	PRO	CD	53.814	11.162	48.256	15.75
1041	GLN	N	50.762	9.495 45.12		
1042	GLN	CA	49.356	9.444 44.80		
1043	GLN	C	48.588	8.192 45.38		
1044	GLN	0	47.377	8.070 45.25		
1045		CB	49.477	9.419 43.30	9 20.0 42.765	32.71
1046	GLN	CD	49.582	10.837 10.778	41.328	36.57
1047	GLN	CD OE1	49.147	10.778	40.486	35.28
1048	GLN	OE1	49.811	10.217	40.400	33.20

				. '	k			.00	
1049	GLN	NE2	47.927	٠,	11.228			1	
1050	LYS	N	49.312		7.229	46.019	) .	20.39	•
1051	LYS	CA	48.578		6.239	46.799	)	19.10	
1052		C	49.447		5.480	47.773	}	16.46	*.
1053	LYS	0	50.649		5.374	47.569	)	16.58	` ;
1054	LYS	СВ	48.004			45.819		22.59	
1055		CG	49.022	٦.		44.999		24.94	
1056		CD	48.247			44.031		30.53	
			46.873			43.585		32.07	
1057	LYS	CE				42.750		40.14	
	LYS	NZ	46.151	***					
1059		N	48.770			48,805		15.88	
1060	GLU	CA	49.282			50.076		15.37	8
1061	GLU	С	50.274			49.993		15.69	
	GLU	0	51.310	• .		50.666		13.68	
1063	GLU	CB.	48.129			50.969		16.72	* .
1064	GLU	CG	47.224	, , , ,	5.504	51.264		14.29	
1065	GLU	CD	46.086		5,726	50.288	}	18.80	
1066	GLU	OE1	46.254		5.591	49.071		21.62	
1067	GLU		44.996		6.034	50.720	)	21.27	*
	GLU		50.001		2.488	49.145	5 , '	18.42	
1069	GLU		50.982			48.868		20.95	
	GLU		52.285			48.121		20.76	90%
1071	GLU		53.234			48.094		20.49	la Time
1072	GLU	СВ	50.407			47.934	1*	20.78	
1072		CG	48.890			47.935		25.93	
			48.253			46.847		25.99	
	GLU	- 7			APPLIES OF MAIN	45.699		28.63	in the same
1075	GLU		48.672						4
	GLU	OE2	47.378			47.165		25.53	
		N	52,277	* . *		47.470		21.38	
. ,	LYS	CA	53.466	,		46.700		21.50	30
1079	LYS	C	54.120			47.250		21.97	
1080	LYS	0	53.847			46.775		22.57	
1081	LYS	CB <sub>.</sub>	53.121			45.211		22.92	
1082	LYS	CG	53,120		2.175	44.507		27.23	
1083	LYS	CD	53.566		1.981	43.052	2	35.59	
1084	LYS	CE '	54.901		2.700	42.698	3	43.39	
1085	LYS	NZ	56.064	· .	2.530	43.615	5	46.48	
1086	GLU	N	54.995	*	4.585	48.246	3	21.26	
1087		CA	55.858			48.687		22.94	
	GLU		56.922			47.618	1	21.77	
	GLU	0	57.224			46.703	2.7	23.33	
	GLU	СВ	56.631			50.033		22.83	
	GLU		56.303	·· · · .		50.740	4 6	28.82	
1091		CD	56.755			49.946		29.12	
	GLU	OE1	57.947			49.726		33.37	
			55.910			49.582		28.44	
1094	GLU	, UE,Z.	JJ.8 10		2.031	<del>-</del> 73.302		20.74	- · · ·

1100       MET       CG       57.088       9.220 45.629       16.56         1101       MET       SD       56.497       10.824       45.225         1102       MET       CE       57.882       11.386       44.232         1103       ILE       N       60.896       7.720 47.557       18.63         1104       ILE       CA       62.108       7.877 48.322       20.60         1105       ILE       C       63.025       8.809 47.514       20.37         1106       ILE       O       63.395       8.568 46.393       23.05         1107       ILE       CB       62.691       6.493 48.661       24.01         1108       ILE       CG1       61.881       5.871 49.811       25.81         1109       ILE       CG2       64.181       6.578 49.065       24.90         1110       ILE       CD1       61.992       4.357 49.909       27.88         1111       PHE       N       63.380       9.944 48.145       19.97	
1112 PHE CA 64.224 10.949 47.503	18.46
1113 PHE C 65.685 10.633 47.905	19.73
1114 PHE O 66.114 10.938 49.001	18.94
1115 PHE CB 63.753 12.370 47.917	16.35
1116 PHE CG 62.290 12.582 47.628	14.87
1117 PHE CD1 61.851 12.931 46.357	13.55
1118 PHE CD2 61.348 12.399 48.607	16.14
1119 PHE CE1 60.511 13.044 46.035	14.72
1120 PHE CE2 60.000 12.519 48.297	14.63
1121 PHE CZ 59.565 12.811 47.004	14.92
1122 GLU N 66.423 9.955 47.009 23.42 1123 GLU CA 67.730 9.355 47.371 27.04	
	26.67
1124 GLU C 68.809 10.381 47.518 1125 GLU O 69.611 10.299 48.440	27.77
1126 GLU CB 68.229 8.253 46.431 32.84	
1127 GLU CG 67.362 6.947 46.395 43.59	
1128 GLU CD 68.029 5.711 45.646 52.36	
1129 GLU OE1 68.362 5.845 44.439 54.69	
1130 GLU OE2 68.199 4.647 46.298 55.07	
1131 ASP N 68.765 11.386 46.636	25.27
1132 ASP CA 69.664 12.544 46.813	24.77
1133 ASP C 69.573 13.322 48.180	24.82
1134 ASP O 70.550 13.880 48.657 1135 ASP CB 69.487 13.507 45.649	26.69 25.87
1135 ASP CB 69.487 13.507 45.649 1136 ASP CG 68.195 14.296 45.612	28.11
1137 ASP OD1 67.142 13.727 45.835	27.95
1138 ASP OD2 68.296 15.481 45.365	30.09
1139 THR N 68.365 13.360 48.766	22.69
1140 THR CA 68.270 13.979 50.054	20.10

1141	THR	С	67.933	13.020	51.178		20.10
1142	THR	0	67.943	13.373	52.347		20.15
1143	THR	CB	67.476	15.268	50.010		18.17
1144	THR	OG1	66.041	14.985	49.980		15.16
1145	THR	CG2	68.214	16.309	49.052		16.50
1146	ASN	N	67.735	11.756	50.863		21.38
1147	ASN	CA	67.628	10.783	51.969		23.28
1148	ASN	С	66.346	10.889	52.86		23.35
1149	ASN	0	66.353	10.887	54.078		24.28
1150	ASN	СВ	68.918	10.815	52.83		27.97
1151	ASN	CG	69.285	9.445 53.414		32.19	
1152	ASN	OD1	68.963	8.394 52.899		34.42	
1153		ND2	70.048	9.471 54.46		31.14	00.74
1154	LEU	N ,	65.246		52.152		22.74
1155	LEU	CA	63.923	11.233	52.687		21.67
1156	LEU	С	62.917	10.253	52.009		20.94
1157	LEU	0	62.978	9.941 50.83		21.39	
1158	LEU	CB	63.586	12.665	52.212		21.84
1159	LEU	CG	63.600	13.787	53.240		23.27
1160	LEU	CD1	64.055	15.113	52.656		17.13
1161	LEU	CD2	64.224	13.472	54.596		21.62
1162	LYS	N	61.974	9.802 52.783	3	21.10	
1163	LYS	CA	60.920	8.934 52.328	3	19.53	
1164	LYS	С	59.609	9.681 52.550	)	18.34	
1165	LYS	0	59.486	10.402	53.520		20.22
1166	LYS	CB .	61.135	7.645 53.18	5	20.94	
1167	LYS	CG	59.973	6.657 53.250	)	22.87	
1168	LYS	CD	60.414	5.282 53.669	9	29.27	•
1169	LYS	CE	59.161	4.429 53.683	3	32.16	
1170	LYS	NZ	59.380	3.130 54.360	)	36.84	
1171	LEU	N	58.659	9.543 51.618	3	16.41	
1172	LEU	CA	57.406	10.345	51.727	7	16.41
1173	LEU	С	56.161	9.462 51.429	9	16.89	
1174	LEU	0	56.112	8.863 50.367	7	16.71	
1175	LEU	CB	57.542	11.526	50.777	7	15.09
1176	LEU	CG	56.271	12.369	50.729	9	13.98
1177	LEU	CD1	56.382	13.211	49.453	3	15.42
1178	LEU	CD2	56.090	13.271	51.990	)	16.75
1179	THR	N	55.183	9.368 52.377	7	17.65	
1180	THR	CA	54.124	8.352 52.362	2.	17.51	
1181	THR	С	52.766	9.036 52.409	9	16.95	
1182	THR	0 '	52.566	9.943 53.206	3	16.10	
1183		СВ	54.424	7.325 53.497		15.94	
1184	THR	OG1	55.871	7.204 53.749		15.73	
1185			54.295	5.897 53.009		17.67	
1186	LEU	N	51.815	8.672 51.504		15.85	

1187 1188 1189 1190 1191 1192 1193 1194 1195 1196 1197 1198 1200 1201 1202	LEU LEU LEU LEU LEU LEU LEU LEU ILE ILE ILE ILE ILE ILE SER	CA C O CB CG1 CD2 N CA C O CB CG1 CG2 CD1 N	50.464 49.849 49.757 49.614 48.211 47.345 48.368 49.561 48.795 47.299 46.786 49.008 50.497 48.030 51.109 46.772		9.240 51.60° 8.492 52.784 7.284 52.720 9.054 50.317 9.745 50.328 9.428 49.092 11.251 9.233 53.881 8.626 54.964 8.487 54.699 7.492 55.168 9.309 56.286 9.177 56.670 8.763 57.336 7.804 56.438 9.561 54.052	1 7 7 3 2 50.477 1 1 1 9 3 6 9	16.51 19.83 20.59 15.58 15.08 13.12 7 18.46 18.30 18.81 19.41 17.44 16.16 17.37 15.26 18.02	15.75
	SER SER SER SER SER GLU GLU	CA C O CB OG N CA C	45.404 45.002 45.731 44.410 44.258 43.804 43.247 41.728	-	9.579 53.498 10.838 11.814 9.580 54.635 10.789 10.869 12.165	} 52.741 52.767	19.40 1 7 21.28 3 7	21.38 22.64 24.51 22.91 25.57 25.11
1211 1212 1213 1214 1215 1216 1217	GLU GLU GLU GLU GLU GLU ASP	O CB CG CD OE1 OE2 N	40.921 43.646 43.357 44.066 44.307 44.430 41.337		11.581 12.377 11.238 11.603 12.793 10.703 13.722	51.933 50.247 49.297 47.994 47.744 47.259 51.692 51.811	3 7 4 4 9	28.12 27.94 32.29 38.30 43.02 41.87 20.31 18.75
1218 1219 1220 1221 1222 1223 1224 1225	ASP ASP ASP ASP ASP ILE	CA C O CB CG OD1 OD2 N	39.977 39.742 40.185 40.000 38.796 37.691 38.979 39.104		14.960 15.833 15.542 16.828 14.403	50.582 50.485 53.115 53.455 52.994 54.204 49.603	2 5 5 5 1 1 1 8	19.30 17.85 21.92 28.13 28.01 33.77 19.47
1226 1227 1228 1229 1230 1231 1232		CA C O CB CG1 CG2 CD1	38.862 37.511 36.486 38.844 40.112 38.613 40.193		15.728 15.086 13.927 13.1 <u>13</u> 14.468	48.297 48.182 48.315 47.248 47.366 45.803 46.229	2 5 3	20.98 22.51 25.15 22.51 22.32 23.73 22.61

1233	LYS	N	37.553	17.021	47.906	21.11
1234	LYS	CA	36.359	17.863	47.866	20.45
1235	LYS	C	36.188	18.316	46.391	20.80
1236	LYS	O	37.064	18.038	45.587	22.26
1237	LYS	CB	36.663	18.921	48.924	22.37
1238	LYS	CG	36.755	18.276	50.320	25.62
1239 1240 1241	LYS LYS	CD CE NZ	35.354 35.212 33.797	18.201 17.510 17.735	50.911 52.277 52.656	29.83 33.05 35.32
1242	THR	N	35.094	18.956	45.972	19.82
1243	THR	CA	34.963	19.315	44.853	21.59
1244	THR	C	35.971	20.151	44.101	20.83
1245	THR	O	36.245	19.955	42.922	22.59
1246	THR	CB	33.646	20.055	44.669	24.13
1247	THR	OG1	33.609	21.266	45.395	24.73
1248	THR	CG2	32.537	19.144	45.139	30.44
1249	TYR	N	36.550	21.088	44.859	17.73
1250	TYR	CA	37.522	21.987	44.234	15.97
1251	TYR	C	38.926	22.037	44.888	15.46
1252	TYR	O	39.790	22.830	44.552	15.86
1253	TYR	CB	36.862	23.362	44.168	12.55
1254	TYR	CG	36.787	24.053	45.510	12.56
1255 1256	TYR TYR TYR	CD1 CD2 CE1	35.748 37.745 35.638	23.758 25.026 24.506	46.416 45.793 47.593	14.70 14.86 15.48
1257 1258 1259	TYR TYR	CE2 CZ	37.676 36.598	25.747 25.497	47.000 47.858	16.69 16.96
1260	TYR	OH	36.474	26.267	48.979	15.71
1261	TYR	N	39.089	21.152	45.889	14.74
1262	TYR	CA	40.333	21.010	46.643	14.03
1263 1264	TYR TYR	C O	40.453 39.469	19.639 18.949	47.315 47.536 47.613	16.12 17.82 12.74
1265 1266 1267	TYR TYR TYR	CB CG CD1	40.575 39.769 40.334	22.195 22.161 21.536	48.905 50.033	15.40 13.77
1268	TYR	CD2	38.479	22.734	48.947	18.58
1269	TYR	CE1	39.560	21.424	51.207	15.81
1270	TYR	CE2	37.715	22.646	50.134	17.84
1271	TYR	CZ	38.266	21.965	51.235	16.69
1272	TYR	OH	37.544	21.811	52.389	18.92
1273	THR	N	41.697	19.226	47.627	14.21
1274	THR	CA	41.956	17.974	48.377	15.42
1275	THR	C	42.951	18.234	49.465	15.44
127 <u>6</u>	THR	O	43.955	18.887	49.248	15.14
1277	THR	CB	42.630	16.919	47.491	16.15
1278	THR	OG1	41.761	16.554	46.440	16.75

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1279	THR	CG2	43.128		15.64	3	48	.159		15.	
1280	VAL	N	42.638	•	17.71	6	50	.607		14.	25
1281	VAL	CA	43.604	•	17.71	6	51	.687		15.	61
1282	VAL	C	44.212		16.32			.849		15.	
		0	43.534		15.32			.815		16.	
1283	VAL	_	_								
1284	VAL	СВ	42.971		18.21			.028		15.	
1285	VAL	CG1	42.210		19.53			.861		17.	
1286	VAL	CG2	44.018	•	18.32	9	54	.147		15.	
1287	ARG	N	45.545		16.29	6	51	.981		15::	23
1288	ARG	CA	46.259		15.03	4	52	.198		12.	90
1289	ARG	C	47.094		15.09			.452		14.	57
	ARG	0	47.669		16.09			.852		14.	
1290										12.	
1291	ARG	CB	47.213		14.68	•		.060			
1292	ARG	CG	46.431		14.66			.771		14.	
1293	ARG	CD	47.328		14.30			.603		17.	
1294	ARG	NE	46.522	•	14.28	8	47	.372		19.	
1295	ARG	CZ	46.343		15.28	6	46	.563		18.	02
1296	ARG	NH1	46.936		16.43		46	.719		20.	75
1297	ARG	NH2	45.562		15.10			.583		22.	30
		N	47.167		13.94			.075		15.	
1298	GLN									15.	
1299	GLN	CA	48.177		13.85			.083			4
1300	GLN	С	49.286		12.98			.578		14.	
1301	GLN	0	49.094		11.91			.000		15,	
1302	GLN	CB	47.425	•	13.29	5	56	.248		19.	99
1303	GLN	CG	48.352		12.98	6 ·	57	.366		24.	39
1304	GLN	CD	47.446		12.29	4	58	.357		33.	29
1305	GLN	OE1	46.290		11.91	7	58	.100		36.	18
1306	GLN	NE2	48.061		12.05			.508		32.	
	LEU	N	50.472		13.51			.813		14.	
1307								.351		16.	
1308	LEU	CA	51.690		12.89						
1309	LEU	С	52.629		12.58			.537		17.	
1310	LEU	Ο.	52.702		13.26			.555		18.	
1311	LEU	CB	52.425	•	13.84	0		.397		16.	
1312	LEU	CG	52.057	•	14.02	9 ,	51	.920		19.	16
1313	LEU	CD1	52.220		15.47	3	51	.494		18.	62
	LEU	CD2	50.780	•	13.40	8	51	.439		19.	45
1315	GLU	N	53.373		11.51			.358		16.	
1316	GLU	CA	54.428		11.31			.300		16.	
								.632		15.	
1317	GLU	C	55.754		11.45						
1318	GLU	0	56.066		10.78			.676		17.	აⴢ
1319	GLU	CB	54.304			56.95			17.38		
1320	GLU	CG	55.477			57.92			23.65		
1321	GLU	CD	55.430			58.76			30.56	٠.	
1322	GLU	OE1	54.547	(	3.475	59.62	26		31.25		
1323	GLU	OE2	56.295		7.734	58.55	3		32.96		
1324	LEU	N	56.529		12.35			.152		17.	81
		• •					- •			•	

1345 1346 1347 1348 1349 1350 1351 1352 1353 1354 1355 1356 1357 1358 1359 1360 1361 1362 1363 1364 1365 1366	LLLLLLLGGGGGGGGGAAAAAAAALLLLLLLLTTTTTTTT	CA COCGCON CCOCGCON CCOCGCOCGCOCGCOCGCOCGCOCGCOCGCOCGCOCGCO	57.907 58.902 59.000 58.084 59.011 59.999 59.611 59.677 60.757 62.183 62.666 61.533 61.368 60.447 62.199 62.873 64.303 65.247 64.793 66.295 67.105 66.668 65.339 65.339 65.339 65.970 67.827 66.080 65.153 65.096 63.794 68.009 69.278 69.130 70.010 69.921 70.079 71.278 67.912 67.627 66.443 66.193	12.759 11.921 13.937 9.597 56.4 8.276 56.3 8.191 57.0 7.146 57.4 7.880 54.9 6.811 54.4 6.740 52.9 6.896 55.0 9.367 57.1 9.474 57.9 9.189 59.4 8.733 60.1 10.888 11.267 11.033 9.502 59.9 9.396 61.3 8.517 61.7	56. 57. 55. 56. 57. 56. 57. 56. 57. 57. 82. 58. 58. 58. 58. 58. 58. 58. 58. 59. 66. 84. 82. 57. 57. 58. 58. 58. 57. 57. 57. 57. 58. 58. 58. 58. 58. 58. 58. 58. 58. 58	617 452 582 31.34 488 412 046 851 30.56 31.30 35.28 35.62 28.95 29.38 27.85 30.00 37.71 41.16 43.17 45.18 739 353 430 43.51 40.76 41.34	19. 17. 19. 20. 17. 24. 25. 24. 26. 28. 29. 27. 30. 28. 31.	54 81 55 65 83 54 84 84 84 84 84 84 84 84 84 84 84 84 84
1367		0	66.193	8.141 62.8	66 -	43.02		
1368_		_CB	67.386	 10.777		009	39.	
1369 1370	THR THR	OG1 CG2	66.206 68.536	11.364 11.758		461 802	42. 37.	

1371	GLN	N e	65.653	8.252 60.722 40.94	
1372	GLN	CA	64.289	7.831 61.000 41.20	
1373	GLN	С	63.388	8.708 61.881 38.70	
1374	GLN	0	62.322	8.304 62.315 38.13	
1375	GLN	CB	64.284	6.380 61.408 45.37	,
1376	GLN	CG	64.898	5.645 60.250 51.51	* 'y
1377	GLN	CD	64.977	4.182 60.571 56.44	1
1378	GLN	OE1	65.018	3.741 61.710 58.04	
1379	GLN	NE2	65.051	3.403 59.497 61.09	
1380	GLU	N	63.785	9.974 62.092 37.29	27.74
1381	GLU	CA	62.727	10.920 62.488	37.74
1382	GLU	С	61.627	11.026 61.422 10.869 60.217	35.73 34.94
1383	GLU	0	61.806	10.869 60.217 12.311 62.903	43.18
1384	GLU	CB	63.277	13.276 63.617	51.01
1385	GLU	CG	62.269 62.855	14.683 63.932	57.11
1386	GLU	CD OE1	64.057	14.771 64.282	59.70
1387 1388	GLU GLU	OE2	62.106	15.688 63.812	59.23
1389	THR	N	60.452	11.227 61.976	33.44
1390	THR	CA	59.253	11.210 61.132	31.67
1391	THR	C	58.364	12.397 61.416	30.34
1392	THR	Ö	58.213	12.814 62.569	31.62
1393	THR	СВ	58.498	9.890 61.287 29.52	Ÿ —
1394	THR	OG1	58.650	9.203 60.065 31.41	
1395	THR	CG2	57.054	9.944 61.745 27.83	
1396	ARG	N	57.809	12.940 60.314	27.40
1397	ARG	CA	56.900	14.072 60.476	22.96
1398	ARG	С	55.656	14.011 59.676	19.60
1399	ARG	0	55.670	13.570 58.544	18.08
1400	ARG	CB	57.517	15.400 60.125	24.02
1401	ARG	CG	58.772	15.656 60.892	27.06
1402	ARG	CD	59.108	17.126 60.821	30.34
1403	ARG	NE	60.299	17.280 61.631	31.08
1404	ARG	CZ	61.259	18.051 61.210	33.27
1405	ARG	NH1.	61.083	18.824 60.138	28.01
1406	ARG	NH2	62.397	17.980 61.904	34.77
1407	GLU	N	54.607	14.560 60.263	19.61
1408	GLU	CA	53.391	14.722 59.470	20.49
1409	GLU	C	53.307	16.068 58.787	18.35
1410	GLU	0	53.451	17.113 59.403	19.50
1411	GLU	CB	52.147	14.576 60.334	23.73
1412	GLU	CG.	50.804	14.805 59.633	30.58 37.37
1413	GLU	CD .	49.709	15.110 60.669 15.928 61.586	42.91
1414 1415	GLU	OE1 OE2	49.895 48.642	15.928 61.586 14.550 60.567	39.95
1415	ILE	NEZ	53.037	15.988 57.502	17.31
1410		IN	JJ.UJ/	10.000	17.01

1417	ILE	CA	52.824	17.183	56.671	15.44
1418	ILE	С	51.428	17.188	56.122	13.03
1419	ILE	0	50.984	16.205	55.574	13.73
1420	ILE	CB	53.864	17.224	55.509	14.77
1421	ILE	CG1	55.351	17.051	55.964	13.99
1422	ILE	CG2	53.605	18.469	54.616	13.81
1423	ILE	CD1	56.011	18.081	56.937	13.39
1424	LEU	Ν	50.739	18.290	56.267	9.92
1425	LEU	CA	49.468	18.427	55.596	10.85
1426	LEU	С	49.617	19.120	54.238	12.45
1427	LEU	0	50.280	20.137	54.100	13.72
1428	LEU	CB	48.604	19.334	56.441	11.52
1429	LEU	CG	47.671	18.670	57.476	17.76
1430	LEU	CD1	47.397	19.618	58.652	14.86
1431	LEU	CD2	48.036	17.234	57.826	15.60
1432	HIS	N .	48.967	18.566	53.225	10.68
1433	HIS	CA	49.032	19.147	51.860	11.55
1434	HIS	C	47.653	19.632	51.485	12.79
1435	HIS	O-	46.691	18.863	51.473	14.70
1436	HIS	CB .	49.509	18.015	50.917	11.04
1437	HIS	CG	49.846	18.414	49.500	9.68
1438	HIS	ND1	49.049	18.115	48.481	10.25
1439	HIS	CD2	50.975	19.042	49.048	10.64
1440	HIS	CE1	49.666	18.552	47.378	12.62
1441	HIS	NE2	50.846	19.126	47.727	11.21
1442	PHE	N	47.569	20.948	51.199	12.23
1443	PHE	CA	46.274	21.521	50.815	10.89
1444	PHE	$C_{n}$	46.347	21.882	49.348	13.34
1445	PHE	Ô	47.133	22.727	48.947	13.55
1446	PHE	CB	45.985	22.804	51.608	10.80
1447	PHE	CG	46.010	22.514	53.082	11.64
1448	PHE	CD1	47.239	22.635	53.771	11.60
1449	PHE	CD2	44.825	22.131	53.751	14.48
.1450	PHE	CE1	47.320	22.342	55.132	12.91
1451	PHE	CE2	44.901	21.852	55.142	15.96
1452	PHE	CZ	46.143	21.936	55.791	13.40
1453	HIS	N	45.551	21.162	48.554	11.63
1454		CA	45.637	21.261	47.102	10.69
1455	HIS	C	44.419	21.873	46.463	11.28
1456	HIS	0	43.339	21.303	46.375	10.79
1457	HIS	CB	45.985	19.871	46.547	11.78
1458	HIS	CG	46.316	19.861	45.068	10.24
1459	HIS	ND1	46.407	20.911	44.219	13.45
1460	HIS	CD2	46.596	18.751	44.321	9.09
1461	HIS	CE1	46.724	20.470	42.970	8.49
1462	HIS	NE2	46.847	19.132	43.028	12.97

					10.007	40.40
1463	TYR	N	44.598	23.139	46.037	10.13
1464	TYR	CA	43.496	23.797	45.360	11.53
1465	TYR	С	43.543	23.415	43.888	12.75
1466	TYR	0	44.543	23.620	43.217	11.94
1467	TYR	СВ	43.699	25.310	45.529	12.91
1468	TYR	CG	42.453	26.186	45.414	14.41
				26.161	44.267	12.62
1469	TYR	CD1	41.617			
1470	TYR	CD2	42.190	27.059	46.496	14.67
1471	TYR	CE1	40.497	27.009	44.203	14.22
1472	TYR	CE2	41.062	27.896	46.437	15.77
1473	TYR	CZ	40.224	27.857	45.292	16.57
1474	TYR	ОН	39.121	28.676	45.247	18.08
1475	THR	N	42.470	22.797	43.411	13.66
1476	THR	CA	42.557	22.223	42.056	14.87
1477	THR	C	41.794	22.989	40.931	16.37
		0	41.788	22.650	39.752	19.74
1478	THR					13.74
1479	THR	CB	42.122	20.752	42.107	
1480	THR	OG1	40.780	20.610	42.619	14.21
1481	THR	CG2	43.091	19.964	42.981	14.13
1482	THR	Ν	41.108	24.086	41.337	17.79
1483	THR	CA	40.223	24.849	40.406	17.71
1484	THR	С	40.545	26.343	40.315	17.22
1485	THR	0	39.717	27.169	39.955	19.49
1486	THR	СВ	38.737	24.756	40.818	20.00
1487	THR	OG1	38.645	25.274	42.134	22.57
1488	THR	CG2	38.165	23.331	40.809	17.86
1489	TRP	N	41.824	26.648	40.637	13.24
1490	TRP	CA	42.325	28.014	40.474	11.42
					39.219	11.23
1491	TRP	С	43.192	28.106		13.88
1492	TRP	0	44.305	27.599	39.146	
1493	TRP	CB	43.175	28.346	41.714	10.59
1494	TRP	CG	43.522	29.827	41.830	10.86
1495	TRP	CD1	43.572	30.834	40.839	10.44
1496	TRP	CD2	43.944	30.486	43.029	11.23
1497	TRP	NE1	43.985	32.030	41.334	11.38
1498	TRP.	CE2	44.223	31.860	42.696	11.59
1499	TRP	CE3	44.121	30.010	44.341	12.73
1500	TRP	CZ2	44.674	32.751	43.694	9.06
1501	TRP	CZ3	44.569	30.912	45.324	11.86
					45.002	9.08
1502	TRP	CH2	44.846	32.248		
1503	PRO	N	42.673	28.737	38.174	11.63
1504	PRO	CA	43.454	28.717	36.926	12.03
1505	PRO	<b>C</b> .	44.752	29.544	36.926	10.91
1506	PRO	0	44.841	30.605	37.525	11.74
1507	PRO	CB	42.440	29.259	35.916	15.45
1508	PRO	CG	41.096	29.392	36.622	16.92

1509 1510 1511	PRO ASP ASP	CD N CA	41.398 45.746 46.936	29.448 29.023 29.819	38.084 36.201 36.018	12.40 9.83 13.50
1512 1513	ASP ASP	C O	46.686 45.875	31.178 31.324	35.371 34.474	15.42 16.15
1514	ASP	СВ	47.954	29.031	35.195	14.19
1515	ASP	CG	49.381	29.475	35.461	17.10
1516	ASP	OD1	49.628	30.416	36.256	16.38
1517 1518	ASP PHE	OD2 N	50.271 47.308	28.847 32.203	34.883 35.958	16.84 15.66
1519	PHE	CA	46.951	33.601	35.630	13.00
1520	PHE	C	45.496	34.007	35.774	13.65
1521	PHE	0	45.036	34.974	35:189	14.13
1522	PHE	CB	47.522	33.989	34.243	15.03
1523	PHE	CG	49.046	33.819	34.221	16.37
1524	PHE	CD1	49.875	34.865	34.737	12.85
1525	PHE	CD2	49.590	32.627	33.669	15.04
1526	PHE	CE1 CE2	51.275 50.986	34.714 32.515	34.723 33.639	10.54 12.82
1527 1528	PHE PHE	CZ	51.798	33.547	34.161	12.62
1529	GLY	N	44.790	33.186	36.600	12.17
1530	GLY	CA	43.402	33.477	36.870 <sup>^</sup>	12.74
1531	GLY	C	43.130	33.699	38.349	13.53
1532	GLY	0	44.023	33.939	39.149	12.80
1533	VAL	N	41.854	33.622	38.691	14.29
1534	VAL	CA	41.352	33.878	40.056	13.36
1535	VAL	С	40.477	32.685	40.533	15.85
1536	VAL	0	39.970	31.920	39.707	16.84
1537	VAL	CB	40.533	35.165	40.144 39.297	12.64 13.96
1538 1539	VAL VAL	CG1	39.246 41.357	35.111 36.379	39.29 <i>1</i> 39.842	12.56
1540	PRO	N	40.310	32.541	41.888	16.09
1541	PRO	CA	39.326	31.553	42.374	14.29
1542	PRO	C	37.904	31.850	41.841	16.70
1543	PRO	0	37.538	32.942	41.394	15.47
1544	PRO	CB	39.425	31.709	43.893	12.24
1545	PRO	CG	40.797	32.289	44.158	11.83
1546	PRO	CD	41.011	33.237	42.981	14.99
1547	GLU	N .	37.091	30.811	41.895	18.37
1548	GLU GLU	CA C	35.721 34.895	30.970 32.053	41.414 42.123	20.99 22.01
1549 1550	GLU	0	34.014	32.699	41.571	24.26
1551	GLU	CB '	34.978	29.614	41.414	22.54
1552	GLU	CG	35.941	28.443	41.162	31.50
1553	GLU	CD	36.522	27.764	42.465	36.66
1554	GLU	OE1	37.244	28.386	43.315	31.28

	*					'
1555	GLU	OE2	36.201	26.562	42.601	36.17
1556	SER	N	35.218	32.225	43.432	20.38
1557	SER	CA ·	34.520	33.218	44.274	17.88
1558	SER	С	35.369	33.535	45.449	15.08
1559	SER	Ö	36.146	32.701	45.885	14.67
1560	SER	CB	33.107	32.739	44.793	16.14
	SER	OG	33.172	31.454	45.419	13.25
1561				,	46.003	17.02
1562	PRO	N	35.198	34.731		
1563	PRO	CA	35.695	34.996	47.370	16.44
1564	PRO	C ·	35.343	33.936	48.410	16.53
1565	PRO	0	36.174	33.495	49.169	15.74
1566	PRO	CB	35.085	36.348	47.744	16.53
1567	PRO	CG	34.912	37.008	46.367	18.72
1568	PRO	CD	34.518	35.878	45.431	16.72
1569	ALA	. N	34.112	33.456	48.396	16.66
1570	ALA	CA	33.729	32.357	49.341	16.12
1571	ALA	С	34.496	31.057	49.241	15.23
1572	ALA	0	34.901	30.462	50.211	15.53
1573	ALA	СВ	32.226	31.970	49.232	15.29
1574	SER	N	34.701	30.625	48.006	15.32
1575	SER	CA	35.478	29.414	47.823	15.79
1576	SER	C	36.944	29.511	48.170	13.45
		0	37.543	28.637	48.804	13.08
1577	SER	•	· ·			17.95
1578	SER	CB	35.222	28.779	46.447	
1579	SER	OG	35.486	29.689	45.407	26.88
1580	PHE	N	37.464	30.720	47.831	13.85
1581	PHE	CA	38.864	31.032	48.219 <sup>-</sup>	12.76
1582	PHE	C	39.021	31.081	49.708	12.10
1583	PHE	,0	39.917	30.465	50.247	14.79
1584	PHE	CB	39.297	32.409	47.672	14.01
1585	PHE	CG	40.660	32.843	48.229	12.89
1586	PHE	CD1	41.853	32.277	47.736	10.70
1587	PHE	CD2	40.701	33.811	49.277	15.33
1588	PHE	CE1	43.092	32.636	48.324	13.65
1589	PHE	CE2	41.935	34.193	49.864	13.88
1590	PHE	CZ	43.111	33.585	49.388	14.38
1591	LEU	N	38.112	31.849	50.367	13.17
1592	LEU	CA	38.075	32.020	51.857	11.98
1593	LEU	C	37.823	30.760	52.651	11.69
1594	LEU	Ö	38.520	30.483	53.608	10.86
				33.104	52.284	11.49
1595	LEU	CB	37.066			9.71
1596	LEU	CG	37.504	34.528	51.922	
1597	LEU	CD1	38.642	35.024	52.815	12.65
_1598_		_CD2_	36.326	35.472	51.928	12.34
1599	ASN	N	36.872	29.960	52.146	13.52
1600	ASN	CA	36.726	28.586	52.660	14.70

1601 1602 1603 1604 1605 1606 1607 1608 1609 1610 1611 1612 1613 1614 1615 1616 1621 1622 1623 1624 1625 1626 1627 1628 1630 1631 1632 1633 1634 1635 1636 1637 1638 1639 1630 1631 1632 1633 1634 1635 1636 1637 1638 1639 1630 1631 1632 1633 1634 1635 1636 1637 1638 1639 1630 1631 1632 1633 1634 1635 1636 1637 1638 1639 1639 1639 1639 1639 1639 1639 1639	A A A A A P P P P P P P P P P L L L L L	C O CB C O D1 N C C O C C C C C C C C C C C C C C C C	38.019 38.453 35.555 35.254 34.760 35.582 38.701 39.997 41.052 41.697 40.490 41.868 43.009 42.033 44.296 43.329 44.460 41.208 42.131 41.892 42.776 42.039 42.908 42.781 44.376 40.589 40.178 40.391 40.855 38.766 38.772 39.357 38.132 39.363 38.121 38.761 40.227 40.694 42.139 42.514	27.745 27.118 27.891 26.492 26.270 25.465 27.785 27.082 27.661 26.908 27.042 26.456 27.294 25.070 26.727 24.515 25.345 29.021 29.689 29.328 29.036 31.214 32.004 33.509 31.595 29.343 28.962 27.509 27.222 29.443 30.884 31.890 31.220 33.222 29.443 30.884 31.890 31.220 33.222 29.443 30.884 31.890 31.220 33.551 26.595 25.200 25.064 24.340	52.629 53.596 51.921 52.493 53.576 51.694 51.452 51.440 52.365 53.084 49.979 49.766 49.586 49.586 49.592 49.407 52.358 53.281 54.775 55.575 53.107 54.144 53.999 54.123 55.118 56.483 56.789 57.866 56.760 57.201 56.414 58.421 56.859 58.855 58.091 55.788 56.007 56.362 57.275	13.68 13.48 15.31 16.04 17.67 15.77 12.27 11.60 11.61 13.15 11.78 14.00 11.64 10.61 12.07 10.24 10.36 11.41 8.72 9.30 11.24 9.97 8.08 13.37 14.13 14.59 17.66 17.85 20.46 17.85 21.97 19.00 13.68 13.76 13.77
1638	LYS	CA	40.694	25.200	56.007	13.68
			•			
1641	LYS	СВ	40.525	24.284	54.793	17.74
1642	LYS	CG	39.416	23.233	54.857	26.69
1643	LYS	CD	39.759	21.835	55.418	32.82
1644	LYS	CE	38.482	20.926	55.564	34.45
1645	LYS	NZ	38.794	19.520	55.920	39.18
1646	VAL	N	43.001	25.790	55.589	13.89

1647 1648 1649 1650 1651 1652 1653 1654 1655 1656 1657 1658 1661 1662 1663 1664 1665 1667 1668 1670 1671 1672 1673 1674 1675 1676 1677 1678 1679 1680 1681 1682	VALUVAL GGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	CA C C C C C C C C C N C N N N C C C C C	44.422 44.678 45.304 45.238 44.967 46.748 44.126 44.218 43.774 44.486 43.366 44.037 43.028 43.712 43.554 42.626 44.379 42.622 42.097 42.975 43.037 40.849 39.696 38.885 38.166 38.978 43.731 44.592 45.788 46.550 45.172 46.254 46.011 47.237 48.494 49.516		25.885 26.442 25.794 26.728 26.260 26.651 27.628 28.196 27.269 27.063 29.447 30.657 31.823 33.077 34.198 34.342 35.143 26.635 25.723 24.566 24.051 25.081 26.051 26.144 27.160 25.220 24.148 22.981 23.113 22.181 22.503 23.396 24.359 24.604 24.856 25.236 24.856		55.972 57.355 58.182 54.971 53.509 55.285 57.609 58.973 60.133 59.011 58.430 58.494 58.799 59.706 58.580 59.706 61.298 60.976 61.298 60.476 61.587 61.732 62.436 60.295 60.481 61.450 61.695 58.735 61.919 62.671 61.883 62.431		11.86 11.86 11.88 12.00 11.88 12.37 12.93 14.48 16.45 17.22 13.67 15.68 14.21 12.03 16.92 16.63 17.05 21.77 30.11 37.62 42.75 39.25 16.50 17.63 20.41 21.90 14.84 13.92 20.72 20.75 22.62 25.75 20.75
1678	SER	OG	46.254		23.396		58.735		13.92
				,	24.604	•	62.671		20.75
				-					
1683	SER	N	48.416		24.672		60.541		20.77
1684 1685	SER SER	CA C	49.659 50.360		24.779 26.137	•	59.772 59.773		20.37 22.30
1686	SER	Ö	51.559	į.	26.233		59.585		22.94
1687	SER	CB	49.485	•	24.400		58.287		17.21
1688	SER	OG	49.026		23.044		58.151		16.17
1689	LEU	N	49.548	-	27.204		59.963		22.44
1690	LEU	_CA	50.024		28.615		59.942	 ٠	25.58
1691 1692	LEU LEU	CO	50.572 50.849		29.143 30.302		61.293 61.553		29.18 34.54
.002			55.546°		JJ.JJE		2		

1693 1694 1695 1696	LEU LEU LEU	CB CG CD1 CD2	48.951 48.548 49.689 47.444	29.616 29.235 29.422 30.047	59.439 58.027 57.098 57.437	23.62 22.66 23.63 25.59 27.88
1697 1698 1699 1700	SER SER SER	N CA C	50.649 50.934 52.365 53.212	28.205 28.588 28.767 28.114	62.203 63.544 63.829 63.230	28.98 29.17 29.28
1701 1702 1703 1704	SER SER PRO PRO	CB OG N CA	50.428 49.070 52.649 54.068	27.503 27.851 29.669 29.983	64.367 64.442 64.766 64.955	29.56 38.82 28.99 28.15
1705 1706 1707 1708	PRO PRO PRO	C O CB CG	54.893 56.103 53.940 52.734	28.912 28.986 31.338 31.150	65.712 65.759 65.624 66.539	27.34 28.43 29.14 28.91
1709 1710 1711 1712	PRO GLU GLU	CD N CA C	51.769 54.222 54.930 55.561	30.350 27.885 26.738 25.818	65.692 66.248 66.757 65.688	28.77 25.89 27.19 26.00 25.89
1713 1714 1715 1716	GLU GLU GLU	O CB CG CD	56.405 54.031 52.691 51.512	24.951 26.017 25.276 26.145	65.946 67.792 67.504 67.001	33.55 41.02 47.74
1717 1718 1719 1720	GLU GLU HIS HIS	OE1 OE2 N CA	51.521 50.550 55.110 55.663	27.382 25.570 26.077 25.415	67.117 66.459 64.437 63.239	49.25 51.04 21.92 18.96
1721 1722 1723 1724	HIS HIS HIS	C O CB CG	56.588 56.466 54.545 53.668	26.312 27.528 25.023 24.065	62.464 62.564 62.270 63.047	17.83 17.07 20.94 20.70
1725 1726 1727 1728	HIS HIS HIS	ND1 CD2 CE1 NE2	52.366 54.077 51.937 52.998	24.242 22.872 23.170 22.336	63.288 63.652 64.033 64.244	22.42 19.89 18.97 19.93
1729 1730 1731 1732	GLY GLY GLY GLY	N CA C	57.430 58.102 57.110 55.911	25.669 26.449 27.069 26.804	61.628 60.582 59.607 59.695	16.31 14.07 12.35 13.46
1733 1734 1735 -1736	PRO PRO PRO	N CA C	57.602 56.641 55.989 56.633	27.933 28.612 27.621 26.687	58.693 57.833 56.834 56.321	11.56 13.06 12.70 13.68
1737 1738	PRO PRO	CB CG	57.455 58.892	29.724 29.277	57.210 57.324	11.82 12.42

1739	PRO	CD	58.963	28.340	58.496	11.92
1740	VAL	N	54.674	27.911	56.610	12.70
1741	VAL	CA	53.984	27.163	55.557	12.91
1742	VAL	C	54.744	27.391	54.248	13.19
1743	VAL	Ö	55.220	28.484	53.926	11.64
1744	VAL	СВ	52.499	27.578	55.507	13.06
1745	VAL	CG1	51.592	26.621	54.745	14.25
	VAL	CG2	52.321	28.992	54.937	13.39
1747	VAL	N	54.874	26.281	53.520	12.59
1748	VAL	CA	55.295	26.380	52.106	11.98
1749	VAL	C	54.101	26.584	51.157	12.38
1750	VAL	0	53.183	25.771	51.097	15.24
1751	VAL	CB	56.061	25.107	51.706	9.51
	VAL	CG1	57.319	24.946	52.599	9.49
1752	VAL	CG2	56.454	25.144	50.213	10.06
1753		N	54.126	27.689	50.434	10.51
1754	VAL	CÅ	53.077	27.963	49.445	10.15
1755	VAL	_		27.938	48.046	11.07
1756	VAL	С	53.712	28.530	47.785	11.12
1757	VAL	0	54.761	29.351	49.698	9.24
1758	VAL	CB	52.435	and the second second	51.143	8.57
1759	VAL	CG1	51.958	29.592	48.689	9.28
1760	VAL	CG2	51.325	29.657	47.133	10.38
1761	HIS	N	53.032	27.235		8.48
1762	HIS	CA	53.504	27.236	45.747	
1763	HIS	С	52.373	27.222	44.762	10.06
1764	HIS	0	51.254	26.782	45.047	9,96
1765	HIS	CB	54.558	26.153	45.458	8.91
1766	HIS	CG	53.991	24.769	45.162	8.64
1767	HIS	ND1	53.572	24.395	43.919	7.44
1768	HIS	CD2	53.803	23.723	46.048	8.51
1769	HIS	CE1	53.098	23.130	43.981	9.19
1770	HIS	NE2	53.243	22.739	45.290	11.46
1771	CYS	N	52.695	27.766	43.579	10.31
1772	CYS	CA	51.889	27.543	42.384	8.62
1773	CYS	С	52.821	26.996	41.318	9.41
1774	CYS	0	53.697	26.240	41.661	9.11
1775	CYS	CB	51.158	28.783	41.966	9.94
1776	CYS	SG	52.104	30.295	42.054	10.94
1777	SER	N	52.652	27.373	40.056	9.15
1778	SER	CA	53.690	26.848	39.157	8.78
1779	SER	С	55.026	27.639	39.274	8.55
1780	SER	0	56.127	27.100	39.390	8.15
1781	SER	CB	53.108	26.847	37.734	7.93
1782	SER	OG	54.103	26.437	36.830	8.26
1783	ĀLĀ	N	54.879	28.997	39.335	8.61
1784	ALA	CA	56.088	29.824	39.571	7.66



1785	ALA	С	56.362	3	0.302		41.022		8.82
1786	ALA	0	57.443		0.767		41.388	. *	11.75
1787	ALA	СВ	55.994		1.024		38.665		7.00
1788	GLY	N	55.332		0.135		41.872		8.66
1789	GLY	CA	55.504		0.616		43.255		8.32
1790	GLY	C	55.306		2.127	•	43.454		9.76
1791	GLY	Ŏ	55.810		2.695		44.413		10.12
1792	ILE	Ň	54.593		2.756		42.493		11.14
1793	ILE	CA	54.394		4.217		42.519		12.07
1794	ILE	C	52.948		4.722	:	42.312		10.76
1795		Ö	52.428		5.499		43.094		12.85
1796	ILE	СВ	55.434		4.998	k.*	41.644		8.68
1797		CG1	55.274		4.717		40.134	,	9.17
1798	ILE	CG2	56.862		4.670		42.093		8.98
1799	ILE	CD1	56.328		5.365		39.216		7.46
1800	GLY	N	52.269		4.172		41.269		10.38
1801	GLY	CA	50.932		4.709		40.970	. •	9.10
1802	GLY	C	49.783		4.383		41.944		10.90
1803	GLY	0	49.349		5.185		42.763		10.20
1804	ARG	N	49.366		3.089		41.810		11.01
1805	ARG	CA	48.418		2.462		42.759	•	10.63
1806	ARG	C	48.954		2.468		44.215		10.18
1807	ARG	0	48.254		2.844		45.134		10.97
1808	ARG	СВ	48.025		1.034		42.275	1 2	9.12
1809	ARG	CG	47.247		1.120		40.967		9.78
1810	ARG	CD	47.007		9.785		40.293		7.58
1811	ARG	NE	48.188		9.323		39.636	* *****	8.89
1812	ARG	CZ	48.221		8.218 · ·		38.919		9.55
1813	ARG	NH1	47.173		7.474		38.805		10.16
1814	ARG	NH2	49.286		7.911		38.225		9.68
1815	SER	N	50.258		2.092		44.358	• •	10.07
1816	SER	CA	50.854		2.120		45.724		8.26
1817	SER	C	50.864		3.513		46.362		10.19
1818	SER	0	50.529		3.638		47.522	,	10.34
1819	SER	СВ	52.294		1.691	* :	45.717		8.89
1820	SER	OG	52.445		0.336		45.240	•	11.05
1821	GLY	N .	51.192		4.586		45.584		10.44
1822	GLY	CA	51.137		5.968		46.103		8.60
1823	GLY	C	49.743		6.409		46.483		9.62
1824	GLY	Ö	49.523		7.100		47.455		11.83
1825	THR	Ň	48.755		5.971	- 1	45.676		11.37
1826	THR	CA	47.324		6.271		45.940		10.02
1827	THR	C	46.841		5.647		47.237		11.10
1828	THR	Ö	46.328		6.337		48.121		9.82
1829	THR	CB	46.392		5.796		44.796		9.53
1830	THR	OG1	46.832		6.337		43.530		10.14
							the second second		



1831	THR	CG2	44.927	36.128	45.095	8.73
1832	PHE	N	47.096	34.326	47.377	9.10
1833	PHE	CA	46.838	33.635	48.636	10.60
1834	PHE	C .	47.384	34.332	49.931	10.58
1835	PHE	Ö	46.702	34.625	50.911	11.24
1836	PHE	CB	47.311	32.168	48.478	8.88
1837	PHE	CG	47.118	31.324	49.741	9.58
1838	PHE	CD1	45.907	30.608	49.935	11.54
1839	PHE	CD2	48.151	31.261	50.713	11.23
1840	PHE	CE1	45.741	29.809	51.090	6.44
1841	PHE	CE2	47.985	30.464	51.879	9.94
1842	PHE	CZ	46.787	29.741	52.023	6.47
1843	CYS	N	48.703	34.583	49.832	10.70
1844		CA ·	49.413	35.190	50.962	10.70
	CYS			36.625	51.272	10.89
1845	CYS	Ç	49.027	1 1	52.423	12.32
1846	CYS	0	48.945	37.006 35.001	50.816	12.32
1847	CYS	CB	50.929	35.091		
1848	CYS	SG	51.574	33.397	50.718	15.41
1849	LEU	N	48.766	37.424	50.215	10.01
1850	LEU	CA	48.287	38.785	50.451	10.37
1851	LEU	C	46.994	38.815	51.257	10.12
1852	LEU	0	46.890	39.510	52.246	9.30
1853	LEU	CB	48.149	39.524	49.125	9.12
1854	LEU	CG	47.758	40.996	49.225	11.23
1855	LEU	CD <sub>1</sub>	47.427	41.612	47.834	11.32
1856	LEU	CD2	48.778	41.823	50.020	11.96
1857	ALA	N	46.015	37.982	50.789	10.91
1858	ALA -	CA	44.724	37.906	51.492	10.80
1859	ALA	С	44.906	37.426	52.955	11.33
1860	ALA	0	44.454	38.026	53.923	10.76
1861	ALA	CB	43.764	36.998	50.732	7.83
1862	ASP	Ν.	45.678	36.373	53.117	11.75
1863	ASP	CA	45.996	35.908	54.497	11.91
1864	ASP	C	46.600	36.952	55.469	11.55
1865	ASP	0	46.156	37.208	56.590	11.95
1866	ASP	CB	46.840	34.628	54.482	10.16
1867	ASP	CG	46.956	34.094	55.912	13.48
1868	ASP	OD1	45.954	33.856	56.609	12.31
1869	ASP	OD2	48.073	33.934	56.360	12.63
1870	THR	N	47.625	37.598	54.924	10.97
1871	THR	CA	48.347	38.561	55.751	11.02
1872	THR	C	47.504	39.800	55.982	11.79
1873	THR	0 .	47.502	40.323	57.088	11.99
1874_			49.692	39.040	55.137	11.07
1875	THR	OG1	50.625	37.985	55.158	11.80
	THR		50.336	40.218	55.881	10.36

1877	CYS	N	46.764		40.247		54.943		10.65
1878	CYS	CA	45.827		41.365		55.230		10.63
1879	CYS	С	44.740		41.106		56.320		12.96
1880	CYS	0	44.486		41.919	•	57.205		13.56
1881	CYS	CB	45.123		41.924		53.999		10.48
1882	CYS	SG	46.329		42.755		52.937		13.42
1883	LEU	N	44.189		39.878	-8-	56.268		10.64
1884	LEU	CA -	43.255	,	39.449	. *	57.316		11.23
1885	LEU	C	43.865	•	39.322		58.731		12.65
1886	LEU	0	43.280		39.757		59.707		14.44
1887	LEU	CB	42.560		38.133		56.873		9.49
1888	LEU	CG '	41.665		38.328		55.653		8.57
1889	LEU	CD1	40.477		39.186		55.978		10.68
1.890	LEU	CD2	41.173		37.002		55.114		11.48
1891	LEU	Ν	45.089		38.753		58.780		13.24
1892	LEU	CA	45.873		38.751		60.039		12.90
1893	LEU	С	46.154		40.116		60.664		13.36
1894	LEU	0	45.961		40.401		61.827		15.00
1895	LEU	CB	47.209		38.118		59.742	:	12.90
1896	LEU	·CG	47.717		37.083		60.724		17.31
1897	LEU	CD1	47.104	•	37.072		62.115		17.90
1898	LEU	CD2	49.240		37.021		60.648		17.91
1899	LEU	Ν.	46.613		41.024		59.811	:	14.06
1900	LEU	CA	46.725	*,	42.436		60.177		15.83
1901	LEU	С	45.461	•	43.107		60.760		15.86
1902	LEU	0	45.460		43.682	,	61.849		15.48
1903	LÉU	CB	47.201		43.234		58.964		15.91
1904	LEU	ĊG	48.624		43.774		58.904	:	18.29
1905	LEU	CD1	49.052		43.573		57.469		20.36
1906	LEU	CD2	49.641		43.305		59.923	-8-	16.23
1907	MET	N	44.375		42.978		59.989		14.06
1908	MET	CA	43.106		43.427		60.510	*	16.10
	MET	<b>C</b> .	42.674	:	42.850		61.894		18.37
1910	MET	0	42.126		43.531		62.759		17.82
1911	MET	CB	42.118		43.137		59.401	1	17.90
1912	MET	CG	40.713		43.550		59.741		22.88
1913	MET	SD	39.601		43.214		58.369		27.93
1914	MET	CE	40.671		43.783		57.014		21.01
1915	ASP	N	42.996		41.548		62.049		17.66
1916	ASP	CA	42.752		40.766		63.290		18.72
1917	ASP	C	43.468		41.301		64.551		20.72
1918	•	O	42.988		41.449		65.670		18.46 15.77
1919	ASP		-43.151		39.310		62.959	•	15.77 15.78
1920	ASP ASP	CG OD1	42.280 41.335		38.294 38.723		63.654 64.270		15.76
1921 1922	ASP	OD1	41.333		37.090		63.595		12.03
1322	<b>TOL</b>	ODZ	72.000		31.030	.*	99.930		12.00

1923	LYS	N	44.725	41.679	64.309	22.67
1924	LYS	CA	45.442	42.083	65.515	26.79
1925	LYS	C	45.030	43.365	66.118	27.91
1926	LYS	Ö	45.173	43.640	67.292	27.25
1927	LYS	СВ	46.939	42.117	65.368	31.63
1928	LYS	CG	47.623	43.011	64.367	36.21
1929	LYS	CD	49.088	42.561	64.344	41.15
1930	LYS	CE	49.122	41.080	63.907	45.83
1931	LYS	NZ	49.950	40.195	64.789	48.52
1932	ARG	N	44.496	44.183	65.244	29.18
1933	ARG	CA	44.107	45.458	65.814	30.25
1934	ARG	C	42.610	45.701	65.816	28.51
1935	ARG	0	42.101	46.694	66.298	29.83
1936	ARG	СВ	44.909	46.490	65.022	35.96
1937	ARG	CG	44.730	46.304	63.509	35.60
1938	ARG	CD	45.670	47.261	62.794	39.72
1939	ARG	NE	47.048	46.827	62.880	44.72
1940	ARG	CZ	47.922	47.107	61.918	49.49
1941	ARG	NH1	47.618	47.655	60.744	46.55
1942	ARG	NH2	49.174	46.847	62.219	54.27
1942	LYS	N	41.920	44.731	65.188	24.03
1943	LYS	CA	40.522	44.893	64.863	22.66
1945	LYS	C	40.322	46.220	64.139	22.19
1946	LYS	Ö	39.223	46.916	64.337	22.54
1947	LYS	СВ	39.649	44.455	66.104	23.45
1948	LYS	CG	39.743	42.957	66.570	20.10
1949	LYS	CD	39.208	41.813	65.634	20.71
1950	LYS	CE	39.159	40.316	66.166	17.94
1951	LYS	NZ	38.787	39.163	65.278	27.72
1952	ASP	N	41.140	46.551	63.222	21.48
1953	ASP	CA	41.082	47.870	62.548	22.85
1954	ASP	C	41.316	47.877	61.015	21.41
1955	ASP	Ö	42.375	48.119	60.444	21.48
1956	ASP	СВ	41.986	48.893	63.285	25.58
1957	ASP	CG	41.991	50.280	62.654	29.64
1958	ASP	OD1	41.113	50.584	61.818	31.77
1959	ASP	OD1	42.919	51.024	62.980	30.41
1960	PRO	N	40.247	47.578	60.311	20.58
1961	PRO	CA	40.371	47.322	58.886	22.04
1962	PRO	C	40.829	48.515	58.129	24.22
1963	PRO	Ö	41.512	48.463	57.118	22.77
1964	PRO	СВ	38.944	47.032	58.454	23.82
1965	PRO	CG .	38.145	46.699	59.709	22.76
1966	PRO	CD	38.893	47.416	60.819	22.17
1967	SER	N	40.411	49.657	58.676	26.49
1968	SER	CA	40.771	50.891	57.983	29.21
	JLI	O/ N	10.77	33.00	37.000	_0 1



1969	SER	С	42.244	51.221	58.015	28.16
1970	SER	0	42.769	51.948	57.184	30.98
1971	SER	СВ	39.918	52.100	58.446	32.36
1972	SER	OG	38.686	52.172	57.658	37.57
1973	SER	N	42.922	50.561	58.956	25.94
1974	SER	CA	44.384	50.605	58.947	23.97
1975	SER	С	45.142	49.677	58.018	23.07
1976	SER	0	46.361	49.640	58.035	23.41
1977	SER	СВ	44.987	50.283	60.317	24.53
1978	SER	OG	44.902	48.866	60.519	26.18
1979	VAL	N	44.421	48.864	57.246	22.45
1980	VAL	CA,	45.137	48.031	56.270	21.20
1981	VAL	C	45.123	48.557	54.850	21.26
1982	VAL	Ó	44.092	48.802	54.231	22.49
1983	VAL	СВ	45.002	46.493	56.466	22.58
1984	VAL	CG1	44.884	45.658	55.190	20.76
1985	VAL	CG2	44.174	46.080	57.682	17.17
1986	ASP	N	46.355	48.833	54.405	19.92
1987	ASP	CA	46.605	49.285	53.043	19.98
1988	ASP	C	47.081	48.123	52.164	17.61
1989	ASP	Ö.	48.232	47.702	52.203	18.51
1990	ASP	СВ	47.639	50.404	53.178	21.39
1991	ASP	CG	47.956	51.130	51.885	25.31
1992	ASP	OD1	47.820	50.542	50.821	23.58
1993	ASP	OD2	48.409	52.287	51.944	32.08
1994	1LE	N	46.134	47.556	51.413	18.00
1995	ILE	CA	46.483	46.335	50.680	16.17
1996	ILE	C	47.626	46.519	49.725	16.20
1997	ILE	Ö	48.485	45.657	49.701	16.71
1998	ILE	СВ	45.244	45.758	49.959	18.69
1999	ILE -	CG1	44.185	45.428	51.013	20.07
2000	ILE	CG2	45.559	44.493	49.137	16.13
2001	ILE	CD1	42.889	44.877	50.407	22.35
2002	LYS	N	47.646	47.644	48.951	15.61
2003	LYS	CA	48.796	47.807	48.039	16.12
2004	LYS	C	50.135	47.931	48.716	15.62
2005	LYS	Ō	51.153	47.394	48.321	15.12
2006	LYS	СВ	48.689	49.055	47.235	18.84
2007	LYS	CG	47.447	49.010	46.334	31.31
2008	LYS	CD	47.091	50.385	45.684	37.73
2009	LYS	CE	47.284	51.463	46.766	43.40
2010	LYS	NZ	46.243	52.480	46.893	47.06
2011	LYS	N	50.099	48.648	49.820	16.19
2012	LYS	CA	51.282	48.662	50.657	17.17
2013	LYS	C	51.745	47.323	51.261	15.81
2014	LYS	Ŏ	52.929	47.020	51.262	13.38
	•	-	<del>-</del>			

2015 2016 2017 2018 2019 2020 2021 2022 2023 2024 2025 2026 2027 2028 2029 2030 2031 2032 2033 2034 2035 2036 2035	LEU LEU LEU LEU LEU LEU	CB CCD CE NCA CCCG CCD1 CCCD1 CCCD1 CCCCCCCCCCCCCCCCC	51.004 52.262 52.080 51.717 52.068 50.779 51.243 51.721 52.682 50.424 50.125 49.369 51.133 51.731 53.154 54.046 50.805 51.267 50.310 51.468 53.356 54.728 55.705	49.698 50.268 51.628 52.861 54.091 46.493 45.153 44.256 43.538 44.394 42.920 45.253 44.416 43.742 44.176 43.392 43.927 43.151 43.372 41.649 45.507 46.026 45.555	51.724 52.260 52.944 52.072 52.860 51.746 52.147 51.029 51.174 53.259 53.057 53.949 49.870 48.691 48.691 48.055 47.497 46.270 45.083 46.603 48.366 48.214 49.289	21.47 26.41 32.75 37.01 41.44 15.38 13.40 12.42 12.19 17.62 13.70 14.26 11.53 12.42 11.91 10.65 12.94 14.51 14.86 12.64 14.10 13.57 11.98
2036 2037 2038 2039 2040 2041 2042 2043 2044	LEU LEU LEU LEU LEU ASP ASP ASP					
2049 2050 2051 2052 2053 2054 2055 2056 2057	ASP ASP MET MET MET MET MET MET ARG ARG	OD1 OD2 N CA C O CB CG	57.469 57.754 55.430 55.755 56.726 57.544 54.494 53.772 52.297 51.819 56.616 57.544	46.507 44.542 42.623 41.284 41.161 40.258 40.507 39.803 39.007 38.502 42.061 42.017	54.030 54.624 50.818 50.344 49.161 49.155 49.955 51.043 50.399 52.033 48.195 47.025	22.88 24.92 13.08 12.59 10.76 9.86 14.18 20.95 25.25 24.15 11.81 14.09

2061	ARG	C	59.018	42.435	47.298	14.19
2062	ARG	0	59.958	42.214	46.558	15.37
2063	ARG	СВ	57.006	42.765	45.786	15.27
2064	ARG	CG	55.506	42.740	45.458	20.41
2065	ARG	CD	54.800	42.082	44.276	22.57
		NE		42.731	43.066	21.00
2066	ARG		55.067			19.36
2067	ARG	CZ	54.603	42.591	41.803	
2068	ARG	NH1	53.481	42.040	41.379	17.30
2069	ARG	NH2	55.419	43.076	40.926	15.47
2070	LYS	N	59.216	42.912	4 <u>8</u> .567	12.82
2071	LYS	CA	60.619	42.962	49.027	11.88
2072	LYS	С	61.320	41.630	49.166 °	11.52
2073	LYS	0	62.519	41.494	49.116	12.87
2074	LYS	CB	60.733	43.761	50.344	10.90
2075	LYS	CG	60.143	45.169	50.220	10.61
2076	LYS	CD	60.078	45.786	51.630	16.89
2077	LYS	CE	59.704	47.286	51.748	18.25
2078	LYS	NZ	59.840	47.665	53.175	18.64
		N	60.530	40.573	49.403	11.46
2079	PHE					10.66
2080	PHE	CA	61.062	39.240	49.651	
2081	PHE	С	60.961	38.264	48.481	10.63
2082	PHE	0	61.718	37.304	48.372	11.08
2083	PHE	CB	60.312	38.582	50.824	10.87
2084	PHE	CG	60.347	39.458	52.036	10.74
2085	PHE	CD1	61.470	39.374	52.899	14.67
2086	PHE	CD2	59.277	40.336	52.321	13.16
2087	PHE	CE1	61.505	40.174	54.080	14.61
2088	PHE	CE2	59.312	41.139	53.481	11.84
2089	PHE	CZ	60.439	41.068	54.334	12.78
2090	ARG	N	59.991	38.503	47.589	11.78
2091	ARG	CA	60.048	37.752	46.323	11.12
2091	ARG	C	59.438	38.615	45.199	10.70
				39.277	45.416	11.00
2093	ARG	0 .	58.427			
2094	ARG	CB	59.361	36.360	46.491	9.05
2095	ARG	CG	59.547	35.389	45.310	8.54
2096	ARG	CD	58.829	34.077	45.618	8.35
2097	ARG	NE	58.807	33.096	44.485	8.45
2098	ARG	CZ	59.812	32.305	44.135	8.44
2099	ARG	NH1	60.967	32.419	44.711	8.74
2100	ARG	NH2	59.670	31.395	43.187	9.54
2101	MET	N	60.064	38.542	43.986	11.10
2102	MET	CA	59.514	39.256	42.833	10.52
2103	MET	C	58.123	38.755	42.342	12.21
2104	MET	Ŏ	57.716	37.596	42.401	11.69
2105	MET	CB	60.515	39.157	41.664	12.40
2106	MET	CG	60.688	37.693	41.161	13.12
2100	IVIL		55.566	07.000	11.101	

2107 2108	MET MET	SD CE	61.708 63.258	37.587 38.266	39.673 40.330	12.49 14.52
2109	GLY	N	57.384	39.697	41.822	11.82
2110	GLY	CA	56.244	39.387	40.947	10.69
2111	GLY	C	55.014	38.778	41.578	11.80
2112	GLY.	0	54.118	38.340	40.868	11.81
2113	LEU	N	54.995	38.855	42.924	11.43
2114	LEU	CA	53.802	38.497	43.681	11.39
2115	LEU	С	52.518	39.174	43.279	12.20 12.24
2116	LEU	0	52.384	40.390	43.261	10.83
2117	LEU	CB	54.050	38.674	45.197	9.47
2118	LEU	CG CD4	55.281	37.955	45.738 45.320	9.47 9.16
2119	LEU	CD1	55.323	36.465 38.148	45.320	10.38
2120	LEU	CD2	55.336 51.588	38.296	47.234	10.56
2121	ILE	N CA	50.396	38.717	42.057	11.17
2122	ILE	CA	50.590	39.082	40.678	. 13.75
2123	ILE	0	51.436	40.016	40.396	13.73
2124 2125	ILE	CB	49.557	39.790	42.868	11.11
2125	ILE	CG1	49.337	39.212	44.233	9.13
2120	ILE	CG2	48.312	40.138	42.008	14.38
2127	ILE	CD1	48.073	39.967	44.918	8.65
2129	GLN	N	50.157	38.291	39.721	13.18
2130	GLN	CA	50.662	38.385	38.342	12.31
2131	GLN	C	49.766	39.112	37.360	14.51
2132	GLN	Ö	50.188	39.490	36.276	14.93
2133	GLN	CB -	51.037	36.985	37.828	12.10
2134	GLN	CG	52.401	36.558	38.336	13.08
2135	GLN	CD	53.427	37.182	37.460	14.33
2136	GLN	OE1	53.477	36.854	36.292	16.45
2137	GLN	NE2	54.248	38.058	38.029	11.97
2138	THR	N	48.519	39.356	37.823	14.47
2139	THR	CA	47.567	40.130	37.057	13.82
2140	THR	C	46.787	41.158	37.866	15.31
2141	THR	Ö	46.656	41.081	39.092	. 15.21
2142	THR	СВ	46.554	39.229	36.350	12.44
2143	THR	OG1	45.588	38.765	37.287	13.84
2144	THR	CG2	47.152	38.042	35.581	12.80
2145	ALA	N	46.231	42.152	37.125	16.12
2146	ALA	CA	45.346	43.159	37.762	17.15
2147	ALA	С	44.035	42.573	38.343	17.06
2148	ALA	0	43.495	43.018	39.351	16.8 <del>4</del>
2149	ÁLA	CB.	44.987	44.299	36.780	15.40
2150	ASP	N	43.567	41.485	37.685	16.67
2151	-ASP	CA	42:388	40.777	38.209	16.05
2152	ASP	С	42.651	39.978	39.460	14.16

0.450		_	44.040	20.000	40.260	13.25
2153	ASP	0	41.818	39.960	40.360	
2154	ASP	CB	41.797	39.833	37.180	16.36
2155	ASP	CG	40.336	39.658	37.485	18.91
2156	ASP	OD1	39.640	40.638	37.769	19.27
2157	ASP	OD2	39.876	38.522	37.453	19.47
2158	GLN	N	43.857	39.397	39.542	12.74
2159	GLN	CA	44.257	38.815	40.853	12.47
2160	GLN	С	44.351	39.857	41.984	13.42
2161	GLN	0	43.906	39.646	43.101	13.68
2162	GLN	СВ	45.579	38.040	40.754	10.94
2163	GLN	CG	45.467	36.777	39.876	10.93
2164	GLN	CD	46.795	36.101	39.634	11.89
2165	GLN	OE1	47.863	36.660	39.815	13.27
2166	GLN	NE2	46.739	34.859	39.129	10.17
2167	LEU	N	44.892	41.066	41.654	15.21
2168	LEU	CA	44.818	42.209	42.611	13.64
2169	LEU	C	43.408	42.567	43.025	14.41
2170	LEU	Ö	43.078	42.676	44.201	15.65
2170	LEU	CB	45.560	43.441	42.074	12.86
			45.681	44.598	43.045	13.02
2172	LEU	CG CD1		45.889	42.389	12.87
2173	LEU	CD1	46.174		44.235	12.32
2174	LEU	CD2	46.540	44.205		13.43
	ARG	N	42.543	42.691	41.995	
2176	ARG	CA	41.161	42.989	42.303	13.38
2177	ARG	C	40.433	41.964	43.135	14.16
2178	ARG	0	39.716	42.292	44.066	14.64
2179	ARG	CB	40.407	43.216	41.022	13.46
2180	ARG	CG	38.925	43.525	41.286	13.29
2181	ARG	CD	38.171	43.714	39.976	16.52
2182	ARG	NE	36.823	44.228	40.247	20.17
2183	ARG	CZ	35.772	43.485	40.354	19.49
2184	ARG	NH1	35.807	42.186	40.332	20.88
2185	ARG	NH2	34.650	44.103	40.484	21.32
2186	PHE	N	40.683	40.692	42.778	13.41
2187	PHE	CA	40.279	39.551	43.612	13.00
2188	PHE	С	40.781	39.564	45.054	12.42
2189	PHE	0	40.006	39.259	45.937	14.84
2190	PHE	CB ·	40.652	38.200	42.969	12.87
2191	PHE	CG	40.024	37.029	43.724	13.08
2192	PHE	CD1	38.719	36.588	43.410	15.82
2193	PHE	CD2	40.732	36.435	44.792	13.48
2194	PHE	CE1	38.069	35.621	44.224	14.88
2194	PHE	CE2	40.074	35.497	45.610	14.71
	PHE	CZ	38.737	35.4 <i>91</i> 35.123	45.347	13.48
2196				39.939	45.279	12.86
	SER	N	42.069			13.82
2198	SER	CA	42.661	40.146	46.627	13.04

2199 2200	SER SER	C O	41.892 41.566	41.095 40.755	47.511 48.633	15.06 15.36
2200	SER	CB	44.037	40.790	46.596	11.50
2202	SER	OG	44.867	39.808	46.005	19.07
2202	TYR	N	41.541	42.275	46.959	14.37
2204	TYR	CA	40.568	43.166	47.642	13.98
2205	TYR	C	39.191	42.565	47.958	14.72
2206	TYR	Ö	38.720	42.678	49.061	16.92
2207	TYR	CB	40.237	44.390	46.809	15.56
2208	TYR	CG	41.257	45.499	46.833	15.10
2209	TYR	CD1	42.504	45.344	46.171	15.01
2210	TYR	CD2	40.898	46.710	47.477	16.69
2211	TYR	CE1	43.433	46.405	46.232	17.26
2212	TYR	CE2	41.786	47.793	47.473	17.79
2213	TYR	CZ	43.052	47.618	46.872	18.91
2214	TYR	OH	43.970	48.651	46.941	22.03
	LEU	N	38.561	41.892	46.996	14.60
2216	LEU	CA	37.297	41.190	47.263	13.80
2217	LEU	C	37.350	40.153	48.419	13.84
2218	LEU	Ö	36.585	40.029	49.351	15.22
2219	LEU	СВ	36.998	40.456	45.966	15.22
2220	LEU	CG	35.889	40.943	45.032	17.91
2221	LEU	CD1	36.181	40.536	43.608	17.36
2222	LEU	CD2	35.469	42.386	45.200	17.38
2223	ALA	N	38.419	39.344	48.343	13.18
2224	ALA	CA	38.629	38.360	49.433	12.65
2225	ALA	C	38.897	38.956	50.826	14.16
2226	ALA	Ö.	38.371	38.463	51.807	14.00
2227	ALA	СВ	39.768	37.370	49.099	13.02
2228	VAL	N	39.721	40.035	50.884	11.78
2229	VAL	CA	39.918	40.769	52.138	12.53
2230	VAL	C	38.651	41.510	52.651	13.82
2231	VAL	Ö	38.291	41.427	53.828	12.86
2232	VAL	СВ	41.111	41.728	52.070	11.66
2233	VAL	CG1	42.380	40.915	51.747	13.24
2234	VAL	CG2	41.316	42.491	53.376	9.92
2235	ILE	Ν	37.962	42.204	51.727	12.29
2236	ILE	CA	36.692	42.836	52.151	13.94
2237	ILE	С	35.643	41.821	52.683	14.15
2238	ILE	0	35.051	42.017	53.733	14.06
2239	ILE	CB	36.150	43.725	50.998	13.50
2240	ILE	CG1	37.132	44.878	50.705	14.91
2241	ILE	CG2	34.736	44.246	51.344	12.74
2242	ILE	CD1	36.837	45.561	49.370	13.99
2243	GLU	N	35.524	40.682	51.982	13.27
2244	GLU	CA	34.623	39.640	52.511	12.79

2245 2246 2247 2248 2249 2250 2251 2252 2253 2254 2255 2256 2257 2258 2259 2260 2261 2262 2263 2264 2265	GLU GLU GLU GLU GLY GLY GLY ALA ALA ALA LYS LYS LYS	C O CB CCD OE2 N CA C O CB	35.054 34.348 34.469 33.577 32.144 31.662 31.437 36.351 36.996 36.903 36.792 36.936 35.555 35.567 36.717 34.522 33.316 33.511 33.007 32.232	38.960 38.838 38.619 37.436 37.802 38.871 37.016 38.579 38.012 38.905 38.436 40.244 41.194 40.936 41.104 42.628 40.415 40.077 39.126 39.309 39.479	53.810 54.812 51.428 51.758 52.097 51.750 52.719 53.807 55.024 56.279 57.408 56.035 57.129 57.966 59.163 56.649 57.318 58.087 59.305 60.412 57.184	14.22 14.98 11.88 12.69 18.61 19.68 20.07 14.10 11.35 12.42 14.09 12.11 13.24 14.50 15.74 12.85 15.43 17.97 18.17 16.72 17.36
	-			;		
2266 2267 2268 2269 2270 2271 2272 2273 2274 2275 2276 2277 2278 2279 2280 2281 2282 2283 2284 2285 2286 2287	LYS LYS LYS PHE PHE PHE PHE PHE PHE ILE ILE ILE ILE	CG CD CE NCA CO CB CCD1 CE2 CZ NCA CCB1 CCG2	31.843 30.798 30.249 29.750 34.298 34.710 35.681 35.418 35.243 35.710 36.999 34.842 37.423 35.280 36.551 36.741 37.667 36.960 37.307 38.752 39.499 39.745	40.457 39.777 40.618 39.613 38.096 37.136 37.783 37.750 35.871 34.853 34.982 33.806 34.059 32.867 33.006 38.443 39.052 40.030 40.294 39.831	56.080 55.206 54.078 53.147 58.987 60.018 61.042 62.227 59.313 60.327 60.906 60.681 61.897 61.642 62.234 60.524 61.474 62.419 63.547 60.687 59.668 61.608	20.89 19.35 21.45 28.58 17.06 17.26 17.65 17.35 16.04 18.17 17.46 18.56 17.26 16.76 17.16 16.95 18.22 19.58 16.65 17.33

0000		004	40.400	07.700	60 100	16.00
2288	ILE	CD1	40.138	37.768	60.180	17.76
2289	MET	N:	35.924	40.676	61.869	17.76
2290	MET	CA	35.293	41.696	62.708	18.59
2291	MET	C	34.125	41.213	63.554	
2292	MET		33.271	42.006	63.936	18.92
2293	MET	CB ·	34.850	42.902	61.856	18.33
2294	MET	CG	36.096	43.579	61.237	18.10
2295	MET	SD	37.293	44.167	62.479	22.98
2296	MET	CE	36.332	45.615	62.928	22.14
2297	GLY	N	34.097	39.878	63.787	17.81
2298	GLY		33.117	39.341	64.764	19.24
2299	GLY	C <sub>.</sub>	31.932	38.467	64.303	18.12
2300	GLY	Ο.	31.162	37.852	65.052	18.31
2301	ASP	N	31.751	38.493	62.977	16.53
2302	ASP	CA	30.662	37.671	62.462	17.00
2303	ASP	С	31.116	36:272	62.108	18.13
2304	ASP	0	31.346	35.891	60.957	19.06
2305	ASP	CB	29.981	38.377	61.275	18.82
2306	ASP	CG	28.755	37.606	60.749	20.22
2307	ASP	OD1	28.315	36.651	61.382	19.51
2308	ASP	OD2	28.234	37.955	59.688	23.56
2309	SER	N	31.276	35.460	63.182	18.73
2310	SER	CA	31.804	34.126	62.883	16.96
2311	SER	C	30.962	33.233	~61.948	18.39
2312	SER	0	31.432	32.300	61.284	18.33
2313	SER	CB .	32.076	33.353	64.164	17.13
2314	SER	OG .	33.211	33.889	64.866	14.25
2315	SER	, <b>N</b> '	29.647	33.584	61.904	17.50
2316	SER	CA	28.726	32.804	61.055	18.74
2317	SER	С	28.994	32.849	59.528	17.75
2318	SER	0	28.665	31.927	58.776	16.68
2319	SER	СВ	27.259	33.169	61.335	19.67
2320	SER	OG	26.885	34.432	60.758	21.88
2321	VAL	N	29.690	33.918	59.107	17.27
2322	VAL	CA	30.098	33.858	57.707	18.22
2323	VAL	С	30.893	32.658	57.299	18.73
2324	VAL	Ō.	30.749	32.254	56.163	19.63
2325	VAL	CB	30.851	35.081	57.155	20.69
2326	VAL	CG1	32.145	35.419	57.906	17.79
2327	VAL	CG2	29.886	36.225	56.907	23.70
2328	GLN	N	31.676	32.100	58.244	18.28
2329	GLN	CA	32.498	30.973	57.851	19.78
2330	GLN	C	31.709	29.809	57.296	23.25
2331	GLN	Ö	31.866	29.325	56.194	25.12
2332	GLN	CB	33.356	30.549	59.027	20.56
2333	GLN	CG	34.279	29.396	58.650	24.84
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2224	CLN	CD	25 222	29.031	59.768	27.26
2334	GLN	CD	35.223		-	31.04
2335	GLN	OE1	35.456	29.725	60.745 59.602	29.42
2336	GLN	NE2	35.798	27.883		
2337	ASP	N	30.649	29.485	58.075	25.31
2338	ASP	CA	29.810	28.447	57.454	27.02
2339	ASP	С	28.987	28.769	56.202	25.79
2340	ASP	0	28.680	27.943	55.345	25.32
2341	ASP	CB	29.058	27.677	58.540	35.78
2342	ASP	CG -	30.060	26.695	59.218	44.92
2343	ASP	OD1	30.644	25.807	58.529	50.04
2344	ASP	OD2	30.268	26.816	60.445	50.59
2345	GLN	N	28.684	30.074	56.081	23.38
2346	GLN	CA	28.196	30.539	54.780	24.32
2347		C	29.105	30.389	53.573	23.16
	GLN	0	28.683	29.998	52.488	21.27
2348		_			54.845	28.00
2349	GLN	CB	27.869	31.970	-	
2350	GLN	CG	26.784	32.237	55.878	35.72
2351	GLN	CD	26.555	33.724	55.747	43.13
2352	GLN	OE1	26.549	34.281	54.641	48.57
2353	GLN	NĘ2	26.403	34.395	56.909	43.82
2354	TRP	N	30.382	30.678	53.819	20.29
2355	TRP	CA	31.346	30.362	52.750	19.31
2356	TRP	С	31.466	28.909	52.431	19.07
2357	TRP	0	31.450	28.527	51.271	19.20
2358	TRP	СВ	32.784	30.779	53.114	19.26
2359	TRP	ĊG	32.909	32.243	53.480	15.65
2360	TRP	CD1	32.161	33.289	52.953	14.94
2361	TRP	CD2	33.865	32.827	54.394	15.85
2362	TRP	NE1	32.572	34.480	53.472	15.54
2363	TRP	CE2	33.623	34.245	54.377	15.86
2364	TRP	CE3	34.912	32.297	55.183	14.86
			34.420	35.102	55.179	16.78
2365	TRP	CZ2				14.37
2366	TRP	CZ3	35.699	33.183	55.951	
2367	TRP	CH2	35.460	34.572	55.969	15.56
2368	LYS	N	31.535	28.079	53.466	21.19
2369	LYS	CA	31.490	26.660	53.154	24.17
2370	LYS	С	30.299	26.154	52.302	25.28
2371	LYS	0	30.419	25.380	51.350	26.15
2372	LYS	CB	31.535	25.905	54.461	26.43
2373	LYS	CG	31.670	24.409	54.259	32.00
2374	LYS	CD	31.629	23.919	55.676	38.87
2375	LYS	CE	31.917	22.442	55.805	45.47
2376	LYS	NZ	31.660	22.084	57.228	51.74
2377	GLU	N	29.117	26.663	52.680	26.22
2378	GLU	CA	27.928	26.358	51.891	27.87
2379	GLU	C	27.975	26.877	50.455	26.53
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2380	GLU	O	27.835	26.148 <sup>7</sup>	49.496	30.42
2381	GLU	СВ	26.659	26.888	52.548	34.80
2382	GLU	CG	26.313	26.343	53.949	47.61
2383	GLU	CD	25.599	24.966	53.934	55.88
2384	GLU	OE1	24.348	24.925	53.838	59.76
2385	GLU	OE2	26.294	23.937	54.064	61.03
2386	LEU	N	28.258	28.163	50.302	24.19
2387	LEU	CA	28.499	28.736	48.972	23.43
2388	LEU	C	29.566	28.108	48.071	24.08
2389	LEU	Ö	29.553	28.130	46.848	22.85
2390	LEU	CB	29.014	30.112	49.163	25.02
2391	LEU	CG	27.980	31.163	49.085	25.39
2392	LEU	CD1	28.441	32.350	49.899	27.03
2393	LEU	CD2	26.614	30.675	49.478	28.40
2394	SER	N	30.563	27.564	48.762	23.22
2395	SER	CA	31.652	26.937	48.040	24.53
2396	SER	C	31.401	25.574	47.438	26.87
2397	SER	0	32.174	25.053	46.642	26.47
2398	SER	CB	32.858	26.864	48.941	23.92
2399	SER	OG	32.879	25.591	49.577	27.33
2400	HIS	N	30.272	24.958	47.865	29.80
2400	HIS	CA	29.907	23.613	47.358	33.13
2401	HIS	C	30.966	22.498	47.442	33.64
	HIS	0	31.210	21.764	46.489	31.30
2403 2404	HIS	CB	29.302	23.669	45.909	36.51
2404	HIS	CG	28.204	24.708	45.783	40.71
	HIS	ND1	28.213	25.693	44.854	43.97
2406	HIS	CD2	27.063	24.893	46.600	42.68
2407	HIS	CE1	27.003	26.502	45.078	43.60
2408	HIS	NE2	26.408	26.006	46.157	42.67
2409	GLU	NEZ N	31.619	22.430	48.620	35.19
2410	GLU	CA	32.799	21.567	48.661	36.70
2411 2412	GLU	CA	32.611	20.063	48.720	38.60
2412	GLU	0	33.525	19.304	48.448	37.53
2413	GLU	СВ	33.697	21.992	49.795	35.82
	GLU	CG	32.956	21.874	51.117	36.20
2415	GLU	CD	33.983	21.924	52.224	38.96
2416		OE1	34.913	22.725	52.123	36.60
2417	GLU			21.139	53.176	41.14
2418	GLU	OE2	33.890	19.658	49.089	42.63
2419	ASP	N	31.402	18.226	49.009	46.40
2420	ASP	CA	31.156	17.554	47.818	47.19
2421	ASP	С	30.772	18.144	46.977	46.38
2422	ASP	O CB	30.066 30.260	17.938	50.394	50.98
2423 2424	ASP ASP	CG	31.161	17.875	51.659	57.85
2424	ASP	OD1	32.063	17.073	51.717	60.50
<b>242</b> 0	ASP	וטט	JZ.UUJ	17.001	O LA EL	55.50

2426	ASP	OD2	30.999	18.696	52.591	61.02
2427	ASP	OXT	31.255	16.438	47.575	48.57
1	TIP3	OH2	60.719	23.664	43.966	20.00
2	TIP3	1H	60.985	23.573	44.873	20.00
3	TIP3	2H	60.658	24.587	43.766	20.00
4	TIP3	OH2	40.411	32.301	35.797	20.00
5	TIP3	1H	40.442	31.973	36.704	20.00
	TIP3	2H	39.543	32.682	35.681	20.00
7	TIP3	OH2	45.842	40.160	69.804	20.00
8	TIP3	1H	46.390	40.196	70.592	20.00
9	TIP3	2H	46.479	40.472	69.181	20.00
10	TIP3	OH2	53.379	29.910	58.076	20.00
11	TIP3	1H	54.092	29.933	58.712	20.00
12	TIP3	2H	53.330	30.805	57.725	20.00
13	TIP3	OH2	65.665	24.233	43.388	20.00
14	TIP3	1H	66.318	23.787	43.947	20.00
15	TIP3	2H	66.089	25.124	43.306	20.00
16	TIP3	OH2	53.559	24.650	58.363	20.00
17	TIP3	1H	54.093	24.160	58.947	20.00
18	TIP3	2H	53.867	25.523	58.315	20.00
19	TIP3	OH2	64.454	48.312	45.244	20.00
20	TIP3	1H	64.267	48.158	46.175	20.00
21	TIP3	2H	63.857	49.038	45.064	20.00
22	TIP3	OH2	65.964	24.398	54.095	20.00
23	TIP3	1H	65.412	24.176	54.850	20.00
24	TIP3	2H	65.297	24.876	53.591	20.00
25	TIP3	OH2	45.682	25.930	65.899	20.00
26	TIP3	1H	46.136	26.039	66.729	20.00
27	TIP3	2H	45.378	26.840	65.851	20.00
28	TIP3	OH2	41.439	40.049	69.937	20.00
29	TIP3	1H	41.192	39.958	70.811	20.00
30	TIP3	2H	40.941	40.831	69.745	20.00
31	TIP3	OH2	44.346	6.948 53.73	1 20.00	
32	TIP3	1H	44.331	7.414 54.57	4 20.00	
33	TIP3	2H	43.897	7.661 53.21	1 20.00	
34	TIP3	OH2	69.712	33.601	44.219	20.00
35	TIP3	1H	69.242	33.254	44.983	20.00
36	TIP3	2H	69.119	34.180	43.748	20.00
37	TIP3	OH2	58.068	40.012	37.522	20.00
38	TIP3	1H	57.605	39.244	37.688	20.00
39	TIP3	2H	57.929	40.200	36.598	20.00
40	TIP3	OH2	38.179	37.107	67.391	20.00
41	TIP3	1H	38.128	37.041	68.351	20.00
42	TIP3	2H	38.342	38.026	67.301	20.00
43	TIP3	OH2	53.580	42.001	38.764	20.00
44	TIP3	1H	53.963	41.667	39.579	20.00

64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79	TIP3 TIP3 TIP3 TIP3 TIP3 TIP3 TIP3 TIP3	2H OH2 1H 2H OH2	53.329 41.144 41.796 41.093 63.542 63.904 63.730 64.565 64.349 64.342 73.801 73.658 73.755 68.073 68.526 68.460 66.813 66.619 67.059 63.757 75.979 75.933 75.587 44.663 45.028 44.876 26.251 26.739 26.454 43.204 42.767 42.761 48.443	42.839 36.980 37.172 37.769 14.613 14.292 15.559 20.375 19.937 21.308 37.858 37.517 38.813 28.759 28.682 29.576 32.500 33.294 32.915 36.906 36.671 37.774 37.116 36.923 37.969 41.751 41.787 42.631 25.219 25.004 26.133 25.421 24.692 26.233 47.491	39.103 35.497 36.184 34.947 42.941 43.762 42.948 61.477 62.304 61.630 38.356 39.241 38.502 56.071 56.924 55.744 36.292 36.779 35.452 54.912 55.795 54.745 45.918 46.791 45.865 34.799 35.676 34.493 59.180 59.969 58.987 36.259 36.714 36.519 57.210	20.00 20.00
76	TIP3	OH2	43.204	25.421	36.259	20.00
						**
80	TIP3	1H	48.360	47.421	58.167	20.00
81	TIP3	2H	48.503	48.418	56.985	20.00
82	TIP3	OH2	61.254	29.798	38.553	20.00
83	TIP3	1H	61.103	29.506	39.441	20.00
84	TIP3	2H 🐇	60.395	30.126	38.301	20.00
85	TIP3	OH2	76.145	33.804	35.273	20.00
86	TIP3	1H	76.871	33.620	35.878	20.00
87	TIP3		76.462	34.565	34.787	20.00
88	TIP3	OH2	55.588	41.658	31.859	20.00
89	TIP3	1H	55.460	41.366	32.756 31.943	20.00
90	TIP3	2H	55.098	42.494	31.843	20.00

91 92 93 94 95 96 97 98 99	TIP3 OH2 TIP3 1H TIP3 2H TIP3 0H2 TIP3 1H TIP3 2H TIP3 OH2 TIP3 1H TIP3 2H	44.487 44.965 43.913 50.573 51.351 50.818 49.953 49.992 49.758	25.053 24.874 25.788 33.176 32.774 33.516 52.982 52.859 53.917	41.030 41.840 41.262 62.732 63.106 61.865 34.749 35.691 34.634	20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00
100 101	TIP3 OH2 TIP3 1H	47.275 48.071	17.995 17.850	39.661 40.167	20.00 20.00
102	TIP3 2H	47.429	18.857	39.243	20.00
103	TIP3 OH2	71.352	29.089	36.878	20.00
104	TIP3 1H	71.322	29.074	37.839	20.00
105	TIP3 2H	71.180	29.979	36.603	20.00
106	TIP3 OH2	29.430	35.859	66.439	20.00
107	TIP3 1H	29.726	35.302	67.151	20.00
108	TIP3 2H	29.413	36.763	66.780	20.00
109	TIP3 OH2	60.271	6.760 44.65		
110	TIP3 1H	60.468	6.649 45.59		
111	TIP3 2H	60.190	7.711 44.59		00.00
112	TIP3 OH2	37.294	40.087	40.715	20.00
113	TIP3 1H	36.898	40.267	41.578	20.00
114	TIP3 2H	37.588	40.958	40.472	20.00
115	TIP3 OH2	43.748	16.614	44.085 44.960	20.00
116	TIP3 1H	44.120	16.513	44.900	20.00
117	TIP3 2H TIP3 OH2	43.925 68.520	17.542 39.888	46.997	20.00
118 119	TIP3 OH2 TIP3 1H	67.991	39.691	47.769	20.00
120	TIP3 1H	67.978	40.491	46.494	20.00
121	TIP3 OH2	58.983	37.779	38.817	20.00
122	TIP3 1H	59.166	37.703	39.753	20.00
123	TIP3 2H	58.959	38.722	38.652	20.00
1	NO H O1	56.508	33.999	33.158	0.00
2	NO H C2	56.195	34.428	34.475	0.00
3	NO HC3	55.272	33.387	34.975	0.00
4	NO HC4	55.005	32.237	34.328	0.00
5	NO H C5	55.802	31.748	.33.139	0.00
6	NO_H C6	57.040	32.661	33.064	0.00
7	NO_H S11	54.303	33.664	36.352	0.00
8 .	NO_H C12	53.738	31.989	. 36.222	0.00
9	NO_H C13	54.015	31.430	35.040	0.00
10	NO_H C14	53.373	30.194		0.00 .
11	NO_H O15	53.544	29.898	33.386	0.00
12		52.655	29.368	35.270	0.00
13	NO_H N17	52.959	31.222	37.208	0.00

14	NO_H C18	52.258	31.692	38.256	0.00
15	NO_H O19	52.471	32.753	38.871	0.00
16	NO_H C20	51.099	30.781	38.736	0.00
.17	NO_H O21	50.031	31.233	39.053	0.00
18 .	NO_H O22	51.286	29.429	38.924	0.00
19	NO_H C23	55.687	35.865	34.517	0.00
20	NO_H N25	56.853	36.772	34.366	0.00
21	NO_H C31	57.312	37.239	33.194	0.00
22	NO_H C32	58.507	38.073	33.403	0.00
23	NO_H C33	58.662	38.076	34.763	0.00
24	NO_H C34	57.674	37.221	35.392	0.00
25	NO_H O35	57.690	36.879	36.554	0.00
26	NO_H O36	56.825	36.836	32.137	0.00
27	NO_H C37	59.400	38.796	32.575	0.00
28	NO_H C38	60.492	39.457	33.192	0.00
29	NO_H C39	60.621	39.444	34.609	0.00
30	NO_H C40	59.698	38.714	35.403	0.00
31	NO_H 044	61.631	40.169	35.326	0.00
32	NO_H C45	61.145	40.731	36.599	0.00

TABLE C

Table of the orthogonal three dimensional coordinates in Ångstroms and B factors (Ų) for Protein Tyrosine Phosphatase 1B complexed with 5-(4-hydroxy-1,3-dioxo-1,3-dihydro-isoindol-2-ylmethyl)-2-(oxalyl-amino)-4,7-dihydro-5H-thieno[2,3-c]pyran-3-carboxylic acid (Example 4).

No	Amin	o acid	X	Υ	Z	В			
1	GLU	N	59.958		70.181		38.145		50.84
2	GLU	CA	58.803		69.268		38.132		51.38
3	GLU	C	58.809		68.319		36.855		49.72
4	GLU	0	59.460		68.640		35.857		49.04
5	GLU	CB	57.591		70.243		38.223		53.79
6	GLU	CG	56.243		69.633		38.610		57.54
7	GLU	CD	56.368		68.679		39.828		62.60
8	GLU	OE1	56.347		69.190		40.946		64.63
9	GĻU	OE2	56.479		67.454		39.647		64.40
10	GLU	HA	58.898		68.637		39.019		20.00
11	GLU	1HB	57.526		70.850		37.323		20.00
12	GLU	2HB	57.818		70.928		39.043		20.00
13	GLU	1HG	55.736	,	69.097		37.823		20.00
14	GLU	2HG	55.552		70.439		38.879		20.00
15	MET	Ν	57.987		67.209		36.871		46.10
16	MET	CA	57.535		66.645		35.550		42.61
17	MET	С	56.699		67.639		34.673		40.49
18	MET	0	56.698		67.589		33.457		38.08
19	MET	CB	56.671		65.360		35.644		41.62
20	MET	CG	55.206		65.625	•	36.082	•	40.45
21	MET	SD	54.276		64.127		36.149		35.84
22	MET	CE	55.232		63.264		37.441	T <sub>A</sub> ,	39.51
23	MET	Н	57.559	,	66.997		37.759		20.00
24	MET	HA	58.440		66.400		35.000		20.00
25	MET	1HB	57.157		64.654	,	36.311		20.00
26	MET	2HB	56.626		64.870		34.671	,	20.00
27	MET	1HG	54.645		66.290		35.422		20.00
28	MET	2HG	55.188		66.074		37.082		20.00
29	MET	1HE	55.510		63.974		38.230		20.00
30	MET	2HE	56.154		62.833		37.052		20.00
31	MET	3HE	54.636		62.486		37.923		20.00
32	GLU	N	55.933		68.506		35.346		40.04
33	GLU	CA	55.048		69.427		34.645		41.99
34	GLU	C	55.841		70.396		33.686		41.64
35	GLU	0	55.416		70.753		32.599	•	42.33
36	GLU	CB	54.205		70.086		35.723		43.82
37	GLU	CG	52.967		70.759		35.105		47.17
38	GLU	CD	51.943		71.122		36.194		49.31
39	GLU.	.OE1	52.37.5		71.686	•	37.179		48.56
40	GLU	OE2	50.736		70.869		36.056		51.19
41	GLU	Н	55.891		68.348		36.331		20.00
42	GLU	HA	54.387		68.813		34.030		20.00

			<b></b> ;	•	425	
43	GLU	1HB	54.780	70.799	36.314	20.00
44	GLU	2HB	53.860	69.333	36.440	20.00
					34.379	20.00
45	GLU	1HG	52.473	70.118		
46	GLU	2HG	53.234	71.688	34.607	20.00
47	LYS	N	57.077	70.711	34.138	40.99
48	LYS	CA	58.115	71.521	33.434	41.50
49	LYS	С	58.730	70.802	32.190	39.57
50	LYS	0	58.673	71.310	31.076	39.12
51	LYS	CB	59.261	71.933	34.428	45.89
52	LYS	CG	58.918	73.024	35.497	51.71
53	LYS	CD	59.986	73.136	36.630	56.13
54	LYS	CE	59.423	73.478	38.036	59.81
55	LYS	NZ	60.210	72.951	39.186	62.35
56	LYS	H	57.200	70.440	35.092	20.00
	LYS	HA	57.601	72.409	33.064	20.00
57					33.861	20.00
58	LYS	1HB	60.122	72.290		
59	LYS	2HB	59.590	71.028	34.944	20.00
60	LYS	1HG	57.935	72.822	35.923	20.00
61	LYS	2HG	58.812	73.990	35.000	20.00
62	LYS	1HD	60.756	73.854	36.348	20.00
63	LYS	2HD	60.497	72.177	36.711	20.00
64	LYS	1HE	58.404	73.085	38.137	20.00
65	LYS	2HE	59.313	74.564	38.135	20.00
66	LYS	1HZ	61.199	73.268	39.140	20.00
67	LYS	2HZ	60.194	71.902	39.141	20.00
68	LYS	3HZ	59.783	73.242	40.090	20.00
69	GLU	N	59.247	69.571	32.450	38.42
70	GLU	CA	59.583	68.656	31.386	37.30
71	GLU	C	58.523	68.608	30.274	34.58
72	GLU	0 .	58.814	68.798	29.094	33.74
73	GLU	CB	59.912	67.281	31.966	40.63
73 74	GLU	CG	60.000	66.235	30.835	46.16
		CD	60.673	64.854	31.065	49.22
75 70	GLU	_				
76	GLU	OE1	60.508	64.217	32.105	50.86
77	GLU	OE2	61.374	64.376	30.167	51.79
78	GLU	Н	59.306	69.279	33.408	20.00
79	GLU	HA	60.487	69.052	30.916	20.00
80	GLU	1HB	59.208	66.970	32.737	20.00
81	GLU	2HB	60.879	67.339	32.462	20.00
82	GLU	1HG	60.482	66.668	29.959	20.00
83	GLU	2HG	58.967	66.050	30.534	20.00
84	PHE	N	57.266	68.373	30.681	31.46
85	PHE	CA	56.238	68.253	29.653	30.49
86	PHE	С	56.102	69.457	28.733	33.05
87	PHE	Ŏ	56.072	69.315	27.517	31.17
88	PHE	СВ	54.928	67.999	30.280	26.00
89	PHE	CG	53.774	67.886	29.306	23.39
90	PHE	CD1	53.774	66.628	28.992	24.75
				69.002	28.777	23.83
91	PHE	CD2	53.136_		28.272	23.65
92	PHE	CE1	52.145	66.498	27.977	23.15
93	PHE	CE2	52.023	68.860 67.500	27. <del>9</del> 77 27.755	23.13
94	PHE	CZ	51.514	67.599	21.133	22.50

					420	
05	PHE	Н	57.145	68.129	31.644	20.00
95						
96	PHE	HA	56.531	67.409	29.021	20.00
97	PHE	1HB	54.687	68.795	30.988	20.00
98	PHE	2HB	54.998	67.080	30.860	20.00
99	PHE	HD1	53.805	65.747	29.338	20.00
					29.012	20.00
100	PHE	HD2	53.488	70.001		
101	PHE	HE1	51.759	65.506	28.091	20.00
102	PHE	HE2	51.544	69.725	27.536	20.00
103	PHE	HZ	50.632	67.461	27.165	20.00
104	GLU	N ·	56.018	70.665	29.336	34.78
					,	36.92
105	GLU	CA	55.897	71.897	28.527	
106	GLU	С	57.122	72.019	27.553	34.88
107	GLU	0	57.053	72.444	26.408	34.89
108	GLU	CB	55.852	73.091	29.484	42.69
109	GLU	CG	54.488	73.612	30.017	51.81
				75.215	30.166	58.22
110	GLU	CD	54.564			
111	GLU	OE1	55,679	75.818	30.045	60.76
112	GLU	OE2	53.497	75.845	30.355	60.88
113	GLU	Н	55.958	70.729	30.335	20.00
114	GLU	HA	54.994	71.824	27.915	20.00
				73.911	28.885	20.00
115	GLU	1HB	56.246			
116	GLU	2HB	56.555	72.973	30.310	20.00
117	GLU	1HG	54.203	73.129	30.954	20.00
118	GLU	2HG	53.684	73.380	29.316	20.00
119	GLN	N	58.282	71.586	28.060	32.17
120	GLN	CA	59.556	71.631	27.323	32.79
						32.59
121	GLN	С	59.615	70.636	26.126	
122	GLN	0	60.173	71.008	25.095	33.38
123	GLN	CB	60.597	71.291	28.375	38.01
124	GLN	CG	62.059	71.020	27.998	46.72
125	GLN	CD	62.634	70.285	29.249	54.60
	GLN	OE1	62.587	69.070	29.367	58.48
126						
127	GLN	NE2	63.129	71,080	30.204	57.65
128	GLN	Н	58.242	71.334	29.034	20.00
129	GLN	HA	59.693	72.651	26.984	20.00
130	GLN	1HB	60.256	70.391	28.865	20.00
131	GLN	2HB	60.549	72.041	29.167	20.00
132	GLN		62.601	71.953	27.837	20.00
		1HG			,	
133	GLN	2HG	62.187	70.379	27.126	20.00
134	GLN	1HE2	63.670	70.582	30.873	20.00
135	GLN	2HE2	62.955	72.054	30.303	20.00
136	ILE	N ·	59.048	69.416	26.317	30.75
137	ILE	CA	58.941	68.342	25.297	29.79
					*	29.61
1.38	ILE	C .	57.992	68.730	24.090	
139	ILE	0	58.254	68.655	22.886	28.92
140	ILE	CB	58.520	66.966	25.824	28.01
141	ILE	CG1	59.648	66.484	26.709	26.65
142	ILE	CG2	58.389	65.988	24.623	24.67
					27.633	30.61
143	ILE	CD1_	59.272	65.414		
144	ILE	Н	58.661	69.264	27.231	20.00
145	ILE	HA	59.984	68.125	25.132	20.00
146	ILE	HB	57.585	67.027	26.380	20.00
		-				

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147	ILE	1HG1	60.041	67.301	27.314	20.00
			60.483	66.163	26.083	20.00
148	ILE					
149	ILE		59.307	65.970	24.039	20.00
150	ILE -		57.582	66.246	23.946	20.00
151	ILE	3HG2	58.243	64.972	24.970	20.00
152	ILE	1HD1	58.868	64.551	27.108	20.00
153	ILE		58.551	65.761	28.375	20.00
154	ILE	3HD1	60.223	65.162	28.094	20.00
					24.608	30.56
155	ASP	N	56.856	69.222		
156	ASP	CA	55.774	69.774	23.845	34.30
157	ASP	С	56.317	70.948	23.013	36.61
158	ASP	0	56.305	70.858	21.794	37.72
159	ASP	СВ	54.623	70.039	24.829	34.23
160	ASP	ĊĠ	53.266	69.515	24.380	34.09
161	ASP	OD1	53.161	68.398	23.871	34.45
						35.24
162	ASP		52.282	70.192	24.584	
163	ASP	Н	56.711	69.214	25.597	20.00
164	ASP	HA	55.509	68.998	23.135	20.00
165	ASP	1HB	54.530	71.096	25.058	20.00
166	ASP	2HB	54.782	69.545	25.769	20.00
167	LYS	N	56.891	71.996	23.669	38.57
168	LYS	CA	57.394	73.143	22.870	40.78
					21.598	40.89
169	LYS	С	58.187	72.746		
170	LYS	0	57.821	73.058	20.475	41.19
171	LYS	CB	58.195	74.152	23.704	44.73
172	LYS	CG	59.737	74.010	23.716	49.46
173	LYS	CD	60.389	74.662	24.948	52.61
174	LYS	CÉ	61.863	74.244	25.156	51.86
175	LYS	NZ	62.098	72.811	24.890	49.92
176	LYS	H	56.748	71.985	24.653	20.00
				73.642	22.511	20.00
1.77	LYS	HA	56.492			
178	LYS	1HB	57.810.	74.129	24.724	20.00
179	LYS	2HB	57.962	75.153	23.339	20.00
180	LYS	1HG	60.175	74.398	22.793	20.00
181	LYS	2HG	59.978	72.965	23.809	20.00
182	LYS	1HD	59.811	74.375	25.827	20.00
183	LYS	2HD	60.301	75.749	24.895	20.00
184	LYS	1HE	62.176	74.480	26.182	20.00
185	LYS	2HE	62.505	74.843	24.502	20.00
						20.00
186	LYS	1HZ	61.912	72.594	23.887	
187	LYS	2HZ	61.421	72.194	25.397	20.00
188	LYS	3HZ	63.067	72.497	25.108	20.00
189	SER	Ν	59.282	72.003	<sup>-</sup> 21.844	39.93
190	SER	CA	60.160	71.674	20.742	41.11
191	SER	C	59.713	70.466	19.823	41.60
192	SER	Ö	60.502	69.985	19.006	44.13
		СВ		71.196	21.527	41.13
193	SER		61.342			
194	SER	OG	60.972	70.326	22.646	43.06
1.95	_SER_		.59.394	71.615	22.757-	20.00
196	SER	HA .	60.414	72.549	20.149	20.00
197	SER	1HB	61.840	72.111	21.923	20.00
198	SER	2HB	62.201	70.871	20.872	20.00

199	SER	HG	60.144	69.739	22.668	0ل.ع
200	GLY	N	58.471	69.970	20.020	39.95
		CA	58.004	68.791	19.312	36.83
201	GLY					
202	GLY	C	58.868	67.488	19.463	35.39
203	GLY	0	59.151	66.807	18.529	37.15
204	GĻY	H	57.818	70.556	20.489	20.00
205	GLY	1HA	57.969	69.018	18.241	20.00
206	GLY	2HA	57.004	68.542	19.656	20,00
207	SER	N	59.300	67.067	20.659	32.66
208	SER	CA	60.096	65.891	20.842	31.67
209	SER	C	59.562	64.556	21.564	29.26
210	SER	Ö	60.362	63.696	21.864	28.04
211	SER	СВ	61.523	66.205	21.262	32.05
					22.072	36.33
212	SER	OG	61.785	67.365		
213	SER	Н	58.976	67.617	21.428	20.00
214	SER	HA	60.276	65.532	19.858	20.00
215	SER	1HB	62.162	66.169	20.334	20.00
216	SER	2HB	61.990	65.313	21.767	20.00
217	SER	HĠ	61.143	68.124	22.244	20.00
218	TRP	N	58.263	64.341	21.705	26.65
219	TRP	CA	57.678	63.094	22.236	21.95
220	TRP	C	58.092	61.836	21.456	21.43
221	TRP	Ŏ	58.398	60.796	21.999	21.86
222	TRP	СВ	56.162	63.309	22.226	22.70
223	TRP	CG	55.712	64.329	23.250	20.91
		CD1	55.145	65.601	23.071	21.20
224	TRP					21.02
225	TRP	CD2	55.791	64.152	24.658	
226	TRP	NE1	54.895	66.197	24.287	21.97
227	TRP		55.270	65.323	25.295	20:57
228	TRP	CE3	56.277	63.109	25.402	18.68
229	TRP	CZ2	55.184	65.387	26.676	20.86
230	TRP	CZ3	56.215	63:188	26.788	16.67
231	TRP	CH2	55.658	64.311	27.429	19.85
232	TRP	Н	57.686	65.143	21.597	20.00
233	TRP	HA	58.058	62.950	23.247	20.00
234	TRP	1HB	55.644	62.373	22.446	20.00
235	TRP	2HB	55.835	63.613	21.232	20.00
236	TRP	HD1	54.914	66.062	22.121	20.00
237	TRP	HE1	54.495	67.092	24.417	20.00
238	TRP	HE3	56.679	62.229	24.906	20.00
				66.296	27.119	20.00
239	TRP	HZ2	54.819			20.00
240	TRP	HZ3	56.622	62.372	27.369	
241	TRP	HH2	55.652	64.334	28.506	20.00
242	ALA	N	58.142	61.898	20.137	21.75
243	ALA :	CA	58.747	60.754	19.397	21.29
244	ALA	С	60.219	60.530		19.48
245	ALA	0	60.614	59.444	20.080	17.73
246	ALA	CB	58.536	60.943	17.876	21.57
247		_ H	-57.792	62.693_	19.657	-20.00
248	ALA	HA	58.226	59.852	19.697	20.00
249	ALA	1HB	59.132	61.760	17.471	20.00
250	ALA	2HB	57.493	61.183	17.683	20.00
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251	ALA	знв	539		60.035		17.340	20.00
252	ALA	N	61.000		61.573		19.777	21.34
253	ALA ALA	CA	62.397 62.640		61.371		20.129 21.579	20.36 20.18
254 255	ALA	C 0	63.307		60.772 59.773		21.765	23.47
256	ALA	СВ	62.920		62.788		20.000	21.69
257	ALA	Н	60.646		62.340		19.255	20.00
258	ALA	HA	62.858		60.692		19.413	20.00
259	ALA	1HB	62.433		63.449		20.701	20.00
260	ALA	2HB	62.763		63.151		18.988	20.00
261	ALA	3HB	63.990		62.798		20.177	20.00
262	ILE	N	61.932		61.425		22.576	20.36
263	ILE	CA	61.790		60.947		23.989	20.21
264	ILE	С	61.398		59.404		24.061	19.21
265	ILE	0	62.077		58.599		24.654	20.20
266	ILE	CB	60.737		61.851		24.792	21.58
267	ILE	CG1	60.968		63.384	•	24.927	25.33
268	ILE	CG2	60.518		61.317		26.196	23.50
269	ILE	CD1	62.410		63.607		25.264	26.28
270	ILE	Н	61.488		62.254		22.255	20.00
271	ILE	HA	62.789		61.058 61.736		24.405 24.267	20.00
272 273	ILE ILE	HB 1HG1	59.791 60.325		63.814		25.692	20.00
273 274	ILE		60.743		63.949	•	24.032	20.00
275	ILE		61.434		61.342		26.788	20.00
276	ILE		60.157		60.290		26.217	20.00
277	ILE		59.791		61.912		26.743	20.00
278	ILE	1HD1			63.217		24.492	20.00
279	ILE	2HD1	62.691		63.122		26.198	20.00
280	ILE	3HD1	62.627		64.673		25.357	20.00
281	TYR	Ν	60.231		59.064		23.403	18.90
282	TYR	CA	59.663		57.728		23.212	16.59
283	TYR	C	60.620		56.761		22.628	19.69
284	TYR	0	60.722		55.641		23.044	21.05
285	TYR	CB	58.346		57.810		22.413	15.48
286	TYR	ÇG CD1	57.722		56.420 55.713		22.275 23.438	15.58 15.71
287 288	TYR TYR	CD1 CD2	57.298 57.586		55.775		21.044	17.07
289	TYR	CE1	56.771		54.396		23.415	15.33
290	TYR	CE2	57.097		54.458		20.983	18.24
291	TYR	CZ	56.694		53.755		22.144	17.83
292	TYR	OH	56.243		52.455		21.973	16.43
293	TYR	Н	59.824	•	59.847		22.928	20.00
294	TYR	HA	59.466		57.370		24.214	20.00
295	TYR	1HB	58.526		58.248		21.425	20.00
296	TYR	2HB	57.651		58.450		22.951	20.00
297	TYR	HD1	57.367		56.218		24.392	20.00
298	TYR	HD2	57.892		56.285		20.136	20.00
299	TYR	HE1	56.386		.54.090	( 1	24.384	 20.00
300	TYR	HE2	56.999		53.996		20.016	20.00
301	TYR	HH	56.824		51.863		22.447	20.00 20.71
302	GLN	N	61.366		57.183		21.632	20.71

					100	
303	GLN	CA	62.5/9	56.353	21.092	25.54
304	GLN	C	63.491	56.092	22.091	21.33
		0	63.866	54.931	22.226	22.52
305	GLN					
306	GLN	CB	62.737	56.837	19.717	29.74
307	GLN	CG	61.480	56.551	18.818	43.77
308	GLN	CD -	61.124	57.664	17.789	50.76
309	GLN	OE1	61.317	58.863	17.906	53.37
310	GLN	NE2	60.647	57.254	16.639	51.41
311	GLN	Н	61.282	58.134	21.330	20.00
312	GLN	HA	61.918	55.375	20.947	20.00
			63.576	56.262	19.335	20.00
313	GLN	1HB				20.00
314	GLN	2HB	63.034	57.885	19.731	
315	GLN	1HG	60.567	56.384	19.385	20.00
316	GLN	2HG	61.648	55.643	18.247	20.00
317	GLN		60.739	<b>58</b> .175	16.252	20.00
318	GLN	2HE2	60.319	56.444	16.170	20.00
319	ASP	N	63.890	57.146	22.820	22.69
320	ASP	CA	64.915	56.979	23.869	23.75
321	ASP	C	64.453	55.995	24.956	21.78
322	ASP	Ö	65.227	55.192	25.428	21.26
323	ASP	СВ	65.301	58.315	24.526	26.73
					23.564	31.02
324	ASP	CG	65.718	59.428		
325	ASP	OD1	66.369	59.105	22.550	31.00
326	ASP	OD2	65.408	60.604	23.837	32.48
327	ASP	,H	63.674	58.089	22.539	20.00
328	ASP	HA	65.794	<b>56.54</b> 5	23.392	20.00
329	ASP	1HB	66.116	58.169	25.233	20.00
330	ASP	2HB	64.466	58.694	25.112	20.00
331	ILE	N	63.136	56.005	25.269	20.63
332	ILE	CA.	62.626	54.892	26.106	19.38
333	ILE	C	62.664	53.512	25.346	19.66
334	ILE	Ö	63.081	52.492	25.913	18.07
335	ILE	СВ	61.192	55.230	26.721	19.75
					27.894	19.61
336	ILE	CG1	61.197	56.210		
337	ILE	CG2	60.570	54.008	27.384	16.65
338	ILE	CD1	59.917	57.001	28.052	21.44
339	ILE	Н	62.684	56.835	24.927	20.00
340	ILE	HA	63.327	54.768	26.932	20.00
341	ILE	HB	60.568	55.595	25.907	20.00
342	ILE	1HG1	61.987	56.937	27.696	20.00
343	ILE	2HG1	61.490	55.748	28.838	20.00
344	ILE		61.179	53.628	28.194	20.00
345	ILE		60.422	53.213	26.655	20.00
346	ILE		59.582	54.249	27.768	20.00
347	ILE		59.114	56.434	28.480	20.00
348	ILE		59.594	57.364	27.080	20.00
					28.668	20.00
349	ILE		60.051	57.880		
350	ARG	N	62.162	53.475	24.066	19.64
351	ARG	CA	62.288	52.251	23.267	_22.20_
352	ARG	C	63.742	51.742	23.370	22.74
353	ARG	0	63.964	50.589	23.705	20.20
354	ARG	CB	61.788	52.370	21.795	23.80

						(	
355	ARG	CG	60.263	52.326	21.416		2ع.28
356	ARG	CD	59.846	51.717	19.966		38.17
357	ARG	NE	58.356	51.345	19.989		47.71
358	ARG	CZ	57.194	51.166	19.228		46.22
359	ARG	NH1	57.118	51.134	17.914		51.10
360	ARG	NH2	56.016	50.979	19.791		39.82
361	ARG	H	61.917	54.354	23.659		20.00
362	ARG	HA	61.656	51.528	23.768		20.00
			62.257	51.527	21.248		20.00
363	ARG	1HB					20.00
364	ARG	2HB	62.258	53.237	21.335		
365	ARG	1HG	59.801	53.297	21.613		20.00
366	ARG	2HG	59.896	51.626	22.154		20.00
367	ARG	1HD	60.387	50.786	19.796		20.00
368	ARG	2HD	60.080	52.412	19.161		20.00
369	ARG	HE	58.008	51.167	20.902		20.00
370	ARG	1HH1	56.227	51.223	17.478		20.00
371	ARG	2HH1	57.905	50.928	17.355		20.00
372	ARG	1HH2	55.262	50.537	19.303		20.00
373	ARG	2HH2	55.845	51.282	20.734		20.00
374	HIS	Ν	64.746	52.610	23.202		20.00
375	HIS	CA	66.074	51.898	23.241		20.00
376	HIS	C	66.658	51.767	24.678		20.00
377	HIS	Ŏ	67.428	50.861	24.973		20.00
378	HIS	СВ	67.188	52.616	22.410		20.00
379	HIS	CG	66.701	53.751	21.532		20.00
380	HIS	ND1	66.063	53.572	20.339		20.00
381	HIS	CD2	66.954	55.115	21.693		20.00
382	HIS	CE1	65.963	54.797	19.797		20.00
		NE2		55.736	20.591		20.00
383	HIS		66.488	53.577	22.922		20.00
384	HIS	H	64.728		22.851		20.00
	HIS	HA	65.925	50.904			20.00
386	HIS	1HB	67.950	 53.016	23.105		20.00
387	HIS	2HB	67.710	51.887	21.787		
388	HIS	HD1	65.792	52.727	19.934		20.00
389	HIS	HD2	67.557	55.565	22.468		20.00
390	HIS	HE1	65.731	54.961	18.764		20.00
391	GLU	N	66.332	52.581	25.699		24.60
392	GLU	CA	66.817	52.241	27.039		23.35
393	GLU	C	66.236	50.854	27.523		20.84
394	GLU	0	66.743	50.234	28.462		20.77
395	GLU	CB	66.563	53.445	27.980		25.78
396	GLU	CG	67.579	54.531	27.671		34.71
397	GLU	CD	67.550	55.857	28.464		41.97
398	GLU	OE1	67.473	55.873	29.688		47.08
399	GLU	OE2	67.723	56.887	27.831		42.03
400	GLU	Н	65.826	53.435	25.569		20.00
401	GLU	НА	67.896	52.124	26.937		20.00
402	GLU	1HB	66.656	53.140	29.019	-	20.00
403	GLU	2HB	65.552	53.848	27.866		20.00
404	GLU	1HG	67.556	 54.790	 26.615		20.00
405	GLU	2HG	68.561	54.116	27.863		20.00
406	ALA	N N	65:138	50.381	26.878		20.22
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407	ALA	CA	64.425	49.305	27.596	10.65
408	ALA	С.	65.171	47.952	27.675	19.60
409	ALA	Ŏ.	66.080	47.612	26.913	21.31
410	ALA	СВ	62.979	49.307	27.198	17.65
411	ALA	Н	64.736	50.905	26.129	20.00
412	ALA	HA	64.411	49.635	28.630	20.00
413	ALA	1HB	62.886	49.086	26.128	20.00
414	ALA	2HB	62.566	50.300	27.380	20.00
415	ALA	3HB	62.406	48.566	27.755	20.00
416	SER	Ν	64.735	47.217	28.738	17.44
417	SER	CA	65.303	45.958	29.138	17.65
418	SER	С	65.080	44.926	28.069	20.44
419	SER	Ö	64.118	44.975	27.295	20.82
420	SER	СВ	64.662	45.552	30.486	16.87
421	SER	OG	64.525	46.634	31.510	17.03
			63.945		29.221	20.00
422	SER	Н		47.581		
423	SER	HA.	66.386	46.074	29.221	20,00
424	SER	1HB	65.308	44.740	30.885	20.00
425	SER	2HB	63.712	44.975	30.299	20.00
426	SER	HG	64.529	47.621	31.322	20.00
427	ASP	Ν	65.958	43.934	28.082	23.65
428	ASP	CA	65.585	42.963	27.136	25.32
429	ASP	С	66.051	41.684	27.689	24.00
430	ASP	Ō	67.238	41.557	27.964	25.47
431	ASP	СВ	66.321	43.342	25.827	30.03
432	ASP	CG	66.013	42.208	24.837	35.94
433	ASP	OD1	64.835	41.789	24.742	37.82
				41.731	24.221	40.50
434	ASP	OD2	66.971		28.494	20.00
435	ASP	Н	66.864	43.946		
436	ASP	HA	64.508	42.853	26.973	20.00
437	ASP	1HB	67.406	43.380	25.994	20.00
438	ASP	2HB	66.013	44.285	25.384	20.00
439	PHE	N	65.113	40.763	27.872	20.26
440	PHE	CA	65.402	39.506	28.474	17.74
441	PHE	С	64.786	38.500	27.583	19.63
442	PHE	0	63.886	38.850	26.811	18.88
443	PHE	CB	64.780	39.359	29.899	15.74
.444	PHE	CG	65.255	40.400	30.868	14.99
445	PHE	CD1	66.433	40.210		14.29
446	PHE	CD2	64.503	41.531	31.095	11.07
447	PHE	CE1	66.871	41.163	32.441	13.28
	PHE	CE2	64.910	42.478	32.012	12.04
448				_		14.86
449	PHE	CZ	66.114	42.290	32.698	
450	PHE	Н	64.258	40.990	27.405	20.00
451	PHE	HA	66.483	39.375	28.481	20.00
452	PHE	1HB	64.945	38.372	30.333	20.00
453	PHE	2HB	63.697	39.445	29.816	20.00
454	PHE	HD1	67.024	39.314	31.395	20.00
455	PHE	HD2	63.562	41.653	30.585	_20.00
456	PHE	HE1	67.800	41.004	32.982	20.00
457	PHE	HE2	64.290	43.328	32.250	20.00
458	PHE	HZ	66.455	43.015	33.431	20.00
	<del>-</del>					

459	PRO	N	65.253	37.196	27.759	.03
460	PRO	CA	64.636	36.040	27.082	17.74
461	PRO	C	63.151	35.749	27.411	17.59
462	PRO	0	62.694	35.796	28.535	18.11
463	PRO	CB	65.582	34.833	27.412	17.29
464	PRO	CG	66.564	35.298	28.571	17.67
465	PRO	CD.	66.467	36.813	28.562	17.97
466	PRO	HA	64.674	36.227	26.012	20.00
467	PRO	1HB	66.190	34.622	26.529	20.00
468	PRO	2HB	65.034	33.928	27.683	20.00
469	PRO	1HG	66.200	34.943	29.539	20.00
			67.592	34.936	28.459	20.00
470	PRO	2HG				
471	PRO	1HD	67.323	37.192	28.014	20.00
472	PRO	2HD	66.501	37.258	29.556	20.00
473	CYS	Ν	62.429	35.432	26.359	16.91
474	CYS.	CA	61.116	34.821	26.476	16.83
475	CYS	C.	61.155	33.441	25.799	17.10
476	CYS	0	60.446	33.139	24.838	15.98
477	CYS	CB	60.141	35.734	25.788	17.42
478	CYS	SG	60.311	37.521	26.025	21.76
479	CYS	H	62.845	35.661	25.483	20.00
480	CYS	HA	60.833	34.706	27.525	20.00
481	CYS	1HB	59.112	35.435	26.009	20.00
					24.709	20.00
482	CYS	2HB	60.247	35.600		
483	CYS	HG	61.234	38.020	25.205	20.00
484	ARG	N	62.055	32.618	26.311	17.15
485	ARG	CA	62.240	31.246	25.729	19.32
486	ARG	С	61.030	30.297	25.783	19.49
487	ARG	0	60.747	29.643	24.808	19.32
488	ARG	CB -	63.463	30.474	26.274	-24.18
489	ARG	CG	64.803	30.648	25.474	34.75
490	ARG	CD	66.027	31.366	26.089	42.80
491		NE	65.822	31.308	27.538	51.13
492	ARG	CZ	66.692	31.456	28.527	51.47
493		NH1	67.986	31.573	28.252	48.86
		NH2		31.494	29.730	48.27
494	ARG		66.143	•	27.211	20.00
495	ARG	Н	62.418	32.881		
496		HA	62.407	31.422	24.671	20.00
497	ARG	1HB	63.253	29.408	26.156	20.00
498	ARĢ	2HB	63.503	30.537	27.360	20.00
499	ARG	1HG	64.620	31.011	24.461	20.00
500	ARG	2HG	65.175	29.633	25.308	20.00
501	ARG	1HD	66.058	32.421	25.820	20.00
502	ARG	2HD	66.979	30.904	25.819	20.00
503	ARG	HE	64.881	31.300	27.880	20.00
504	ARG		68.672	31.709	28.970	20.00
505	ARG		68.274	31.511	27.301	20.00
506			66.703	31.481	30.549	20.00
			_65.138	31.401 -31.551	29.794	20.00
-507	-ARG			-3.1.55.1 30.155	26.945	17.97
508		N	60.319			
509	VAL	CA	59.174	29.238	26.975	16.80
510	VAL	С	58.113	29.793	26.009 <sup>-</sup>	15.05

511	VAL	0	57.407	29.009	25.329	155
512	VAL	CB ·	58.828	28.750	28.462	18.01
513	VAL	CG1	57.372	28.681	28.911	15.99
514	VAL	CG2	59.711	29.222	29.601	17.07
			60.614	30.724	27.716	20.00
515	VAL	Н				
516	VAL	HA	59.553	28.350	26.498	20.00
517	VAL	НВ	59.085	27.690	28.412	20.00
518	VAL		56.744	28.248	28.134	20.00
519	VAL	2HG1	56.988	29.671	29.154	20.00
520	VAL	3HG1	57.249	28.037	29.790	20.00
521	VAL	1HG2	59.485	30.243	29.907	20.00
522	VAL		60.768	29.174	29.338	20.00
523	VAL		59.588	28.576	30.471	20.00
524	ALA	N	57.929	31.159	25.879	14.53
		CA	56.893	31.736	24.965	14.43
525	ALA				23.579	15.46
526	ALA	C	57.034	31.215		
527	ALA	0	56.026	30.897	22.995	16.10
528	ALA	CB	56.950	33.283	24.793	12.61
529	ALA	Н	58.489	31.728	26.477	20.00
530	ALA	. HA	55.936	31.377	25.317	20.00
531	ALA	1HB	57.978	33.586	24.605	20.00
532	ALA	2HB	56.694	33.786	25.717	20.00
533	ALA	ЗНВ	56.439	33.685	23.923	20.00
534	LYS	N	58.297	31.191	23.122	17.52
535	LYS	CA	58.835	30.716	21.906	17.70
536	LYS	C	58.993	29.228	21.723	19.60
		Ö	59.486	28.823	20.702	22.60
537	LYS					19.24
538	LYS	CB	60.168	31.413	21.780	
539	LYS	CG	60.083	32.927	21.737	19.97
540	LYS	CD	59.064	33.361	20.674	21.69
541	LYS	CE	59.193	34.787	20.078	24.74
542	LYS	NZ	58.108	35.143	19.082	26.46
543	LYS	Н	58.962	31.547	23.782	20.00
544	LYS	HA	58.145	31.021	21.117	20.00
545	LYS	1HB	60.687	31.055	20.890	20.00
546	LYS	2HB	60.838	31.130	22.595	20.00
547	LYS	1HG		33.367	21.526	20.00
548	LYS	2HG	59.740	33.340	22.686	20.00
549	LYS	1HD	58.049	33.228	21.050	20.00
				32.677	19.824	20.00
550	LYS	2HD	59.142			
551	LYS	1HE	60.168	34.871	19.582	20.00
552	LYS	2HE	59.209	35.527	20.883	20.00
553	LYS	1HZ	57,167	35.181	19.539	20.00
5 <b>5</b> 4	LYS	2HZ	58.096	34.490	18.275	20.00
555	LYS	3HZ	58.222	36.119	18.723	20.00
556	LEU	Ν	58.600	28.321	22.639	20.39
557	LEU	CA	58.649	26.874	22.415	19.13
558	LEU	C	57.499	26.495	21.561	21.96
559	LEU	Ö	56.401	27.065	21.641	20.82
560	LEU	CB	58.382	26.108	23.763	18.89
561	LEU	CG	59.526	26.182	24.733	17.87
	LEU	CD1	60.698	25.466	24.168	19.14
562	LEU	CDI	00.030	20.400	24.100	10.17

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563	LEU	CD2	59.172	25.541		26.090		17.	39
564	LEU	Н	58.246	28.683		23.495		20.	00
565	LEU	HA	59.615	26.655		21.958		20.	00
566	LEU	1HB	58.175	25.056		23.593		20.	00
567	LEU	2HB	57.485	26:522		24.240	<i>?</i>	20.	00
568	LEU	HG	59.806	27.235		24.855		20.	00
569	LEU	1HD1	61,105	25.936		23.272		20.	00
570	LEU	2HD1	60.454	24.433		23.933		20.	00
571	LEU	3HD1	61.510	25.440		24.897		20.	00
572	LEU	1HD2	58.924	24.482		25.978		20.	00
573	LEU	2HD2	58.292	26.034		26.506		20.	00
<sub>-</sub> 574	LEU	3HD2	59.988	25.638		26.811		20.	00
575	PRO	N	57.678	25.487		20.700		23.	61
576	PRO	CA	56.624	25.246		19.703		24.	44
577	PRO	С	55.294	24.893	•	20.278	,	22.	23
578	PRO	0	54.301	25.224		19.668		23.	82
579	PRO	CB	57.127	24.113		18.808		26.	16
580	PRO	CG	58.632	24.213		19.010		<b>27</b> .	75
581	PRO	CD	58.950	24.825		20.348		26.	72
582	PRO	HA	56.514	26.154		19.103		20.	00
583	PRO	1HB	56.822	24.188		17.756		20.	00
584	PRO	2HB	56.800	23.128		19.167		20.	00
585	PRO	1HG	59.145	23.261		18.859		20.	00
586	PRO	2HG	59.017	24.887		18.240		20.0	00
587	PRO	1HD	59.763	25.549	. *	20.239		20.0	0Q
588	PRO	2HD	59.256	24.094		21.089		20.	00
589	LYS	N	55.275	24.280		21.486		22.	30
590	LYS	CA	53.990	23.915		22.172		22.2	26
591	LYS	С	·53.174	25.094		22.666		22.9	91
592	LYS	0 :	51.958	24.974		22.806		25.9	95
593	LYS	CB	54.084	22.876		23.305		23.	13
594	LYS	CG	54.925	23.304		24.549		24.	19
595	LYS	CD	54.938	22.261		25.694		27.4	
596	LYS	CE	55.785	22.825		26.873		31.6	<del>6</del> 5
597	LYS	NZ	55.815	22.067		28.141		37.	
598	LYS	Н	56.127	23.877		21.799		20.0	
599	LYS	HA	53.394	23.472		21.378		20.0	
600	LYS	1HB	54.522	21.963		22.895		20.0	
601	LYS	2HB	53.080	22.604		23.629		20.0	
602	LYS	1HG	54.556	24.257		24.934		20,0	
603	LYS	2HG	55.948	23.498		24.234		20.0	
604	LYS	1HD	55.299	21.288		25.356		20.0	
605	LYS	2HD	53.903	22.109		26.017		20.0	
606	LYS	1HE	55.373	23.814		27.106		20.0	
607	LYS	2HE	56.810	 23.019		26.530		20.0	
608	LYS	1HZ	56.128	21.082		28.050		20.0	
609	LYS	2HZ	54.827	22.030		28.497		20.0	
610	LYS	3HZ	56.381	22.529		28.880		20.0	
611 <sup>-</sup>	ASN		53.890	 26.227		22.879		20.4	
612	ASN	CA	53.214	27.434		23.253		18.9	
613	ASN	C	52.746	28.350		22:087		19.8	
614	ASN	0	52.113	29.392		22.300		16.6	51

615	ASN	CD	E		20 440		24.400		15 00
616	ASN	CB CG	54.191 54.146		28.118 27.319		24.160 25.455		15.32 18.08
617	ASN	OD1	53.155	1.1	26.658	٠.,	25.743		16.89
618	ASN	ND2	55.231		27.426		26.237		17.04
619	ASN	Н	54.866		26.232		22.663		20.00
620	ASN	HA	52.284		27.169		23.748		20.00
621	ASN	1HB	53.876	. '	29.132		24.406		20.00
622	ASN	2HB	55.207		28.155	•.	23.763	. •	20.00
623 624	ASN ASN	1HD2 2HD2	2 55.118 2 56.036		26.972 27.928		27.105 25.914		20.00 20.00
625	LYS	N ,	53.052		27.988		20.818		19.50
626	LYS	CA	52.880		28.993		19.736		20.65
627	LYS	C ·	51.462		29.515		19.709		17.99
628	LYS	0	51.176	,	30.690		19.685		17.79
629	LYS	СВ	53.224		28.321		18.344	٠.	24.77
630	LYS	CG	53.732		29.255		17.235		35.23
631 632	LYS LYS	CD CE	54.019 54.433		28.612 29.635		15.850 14.731		43.17 48.79
633	LYS	NZ	54.133		29.061		13.411		51.89
634	LYS	Н	53.580		27.148	•.	20.703		20.00
635	LYS	НА	53.557		29.817		19.962		20.00
636	LYS	1HB	52.351		27.791		17.957		20.00
637	LYS	2HB	53.960		27.534		18.498		20.00
638 639	LYS LYS	1HG 2HG	54.655		29.727		17.587		20.00
640	LYS	1HD	53.003 53.098		30.061 28.115	٠.	17.104 15.541	•	20.00
641	LYS	2HD	54.759		27.817		15.949		20.00
642	LYS	1HE	55.496		29.890		14.790		20.00
643	LYS	2HE	53.870		30.569		14.815		20.00
644	LYS	1HZ	53.097		29.011		13.287		20.00
645 646	LYS LYS	2HZ 3HZ	54.453 54.503		28.080 29.603		13.339 12.605		20.00 20.00
647	ASN	N	50.566		28.527		19.762		16.87
648	ASN	CA	49.107		28.843		19.726		17.37
649	ASN	C	48.464		29.507		21.029		14.52
650	ASN	0	47.248		29.727	• (	21.164		14.87
651 652	ASN	CB	48.373	٠	27.533	•	19.306		17.56
653 ·	ASN ASN	CG OD1	48.272 48.877		26.488 26.514		20.402 21.451		21.16 23.28
654	ASN	ND2	47.525		25.472		20.024		22.37
655	ASN	. Н	50.925		27.620		19.959	•	20.00
656	ASN	HA ,	48.961		29.568		18.921	•	20.00
657	ASN	1HB	48.842		27.093		18.437		20.00
658 659	ASN ASN	2HB	47.350 47.520		27.783 24.710		19.013		20.00
660	ASN		47.013		25.510		20.667 19.175		20.00
661	ARG	N	49.398		29.743		21.971		13.45
662	ARG	CA	49.080		30.421		23.219		14.08
663	ARG	C	49.442		31.897		23.164	• 6.	11.72
664 665	ARG	O	49.239		32.606		24.120	- 1	11.90
665 666	ARG ARG	CB CG	49.812 49.139		29.746 28.440		24.405 24.752		14.72 15.03
555	AING		TU. 100		20.440		24.1 JZ		13.03

ARG	CD ·	49 763		27 746		25 988		13.22
								14.96
								15.87
ARG	NH1	49.281		26.474		28.507		14.76
ARG	NH2	48.299		24.741		27.436		16.81
								20.00
								20.00
								20.00
ARG	2HB	50.877		29.631		24.213		20.00
ARG	1HG	49.197		27.772		23.893		20.00
								20.00
								20.00
								20.00
ARG	HE	48.517		26.130		25.480		20.00
ARG	1HH1	49.168		26.032		29.395		20.00
								20.00
								20.00
								20.00
ASN	Ν	49.996		32.318		22.035		12.70 ·
ASN	CA	50.406		33.706		21.870		12.44
	C	49 508				20.783		12.58
								11.79
								15.34
								15.14
ASN	OD1	52.540		33.681		23.718		13.40
ASN	ND2	53.664		32.406		22.420		15.01
								20.00
								20.00
								20.00
AŞN	2HB	52.099		33.162				20.00
ASN	1HD2	54.227		32.167	/	23.194		20.00
ASN	2HD2	53.844		32.004		21.533		20.00
								11.31
								10.21
	• .				•			
								11.77
ARG	0	48.660		36.681°		17.785		10.43
ARG	CB	47.337		37.044		20.779		10.53
ARG	CG	46.560		37.866		19.869		9.14
								10.01
								10.47
								10,78
ARG	NH1	43.385		38.316		19.382		11.53
ARG	NH2	42.642		37.281		21.300		10.00
	Н	48 857		35 623		22 040	•	20.00
								20.00
								20.00
								20.00
ARG	1HG	45.916		37.234		19.260		20.00
-ARG-	2HG	47.204	-p =	38.380		19.156		20.00
								20.00
	2HD	46.466		39.497		21.205		20.00
ARC:	/ [ ]							
ARG ARG	HE	44.748		38.186		22.399		20.00
	ASN ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	ARG	ARG NE 48.959 ARG CZ 48.858 ARG NH1 49.281 ARG NH2 48.299 ARG H 50.337 ARG HA 48.004 ARG 1HB 49.727 ARG 2HB 50.877 ARG 1HG 49.197 ARG 2HG 48.078 ARG 1HD 49.685 ARG 2HD 50.798 ARG 1HH 49.168 ARG 1HH 49.168 ARG 2HH1 49.705 ARG 1HH2 48.173 ARG 2HH2 47.960 ASN N 49.996 ASN CA 50.406 ASN CA 50.406 ASN CB 51.893 ASN CG 52.706 ASN ND2 53.664 ASN ND2 53.664 ASN H 50.101 ASN HA 50.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.311 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.311 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 48.957 ARG CA 48.660 ARG CB 47.337 ARG CG 46.560 ARG CD 45.810 ARG HA 47.516 ARG HA 47.516 ARG 1HB 47.990 ARG 2HB 46.686 ARG 1HB 47.990 ARG 2HB 46.686 ARG 1HG 45.916 -ARG 2HG 47.204	ARG NE 48.959 ARG CZ 48.858 ARG NH1 49.281 ARG NH2 48.299 ARG H 50.337 ARG HA 48.004 ARG 1HB 49.727 ARG 2HB 50.877 ARG 1HG 49.197 ARG 2HG 48.078 ARG 1HD 49.685 ARG 1HD 49.685 ARG 2HD 50.798 ARG 1HH1 49.168 ARG 2HH1 49.705 ARG 1HH2 48.173 ARG 2HH2 47.960 ASN N 49.996 ASN CA 50.406 ASN C 49.508 ASN O 49.360 ASN CB 51.893 ASN CG 52.706 ASN OD1 52.540 ASN ND2 53.664 ASN H 50.101 ASN HA 50.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.211 ASN 2HB 52.099 ASN 1HD2 54.227 ASN 1HB 52.3844 ARG N 48.922 ARG CA 48.157 ARG C 49.030 ARG O 48.660 ARG CB 47.337 ARG CG 46.560 ARG O 48.660	ARG NE 48.959 26.521 ARG CZ 48.858 25.909 ARG NH1 49.281 26.474 ARG NH2 48.299 24.741 ARG H 50.337 29.472 ARG HA 48.004 30.372 ARG 1HB 49.727 30.401 ARG 2HB 50.877 29.631 ARG 1HG 49.197 27.772 ARG 2HG 48.078 28.628 ARG 1HD 49.685 28.396 ARG 2HD 50.798 27.458 ARG 1HH 49.168 26.032 ARG 1HH1 49.168 26.032 ARG 1HH2 48.173 24.251 ARG 2HH2 47.960 24.316 ASN N 49.996 32.318 ASN CA 50.406 33.706 ASN C 49.508 34.267 ASN O 49.360 33.682 ASN CB 51.893 33.770 ASN CG 52.706 33.250 ASN OD1 52.540 33.681 ASN ND2 53.664 32.406 ASN H 50.101 31.681 ASN HA 50.227 34.257 ASN 1HB 52.211 34.789 ASN 2HB 52.099 33.162 ASN 1HD2 54.227 32.167 ASN 2HB 52.099 33.162 ASN 1HD2 54.227 32.167 ASN 2HD2 53.844 32.004 ARG N 48.922 35.407 ARG CA 48.157 36.050 ARG CB 47.337 37.044 ARG CB 47.337 37.044 ARG CG 46.560 37.866 ARG CD 45.810 38.876 ARG CB 47.337 37.044 ARG CA 48.644 38.387 ARG CA 48.660 36.681 ARG CB 47.337 37.044 ARG NH1 43.385 38.316 ARG NH2 42.642 37.281 ARG H 48.857 35.623 ARG HA 47.516 35.294 ARG 1HB 47.990 37.689 ARG 2HB 46.686 36.526 ARG 1HG 45.916 37.234	ARG NE 48.959 26.521 ARG CZ 48.858 25.909 ARG NH1 49.281 26.474 ARG NH2 48.299 24.741 ARG H 50.337 29.472 ARG HA 48.004 30.372 ARG 1HB 49.727 30.401 ARG 2HB 50.877 29.631 ARG 1HG 49.197 27.772 ARG 2HG 48.078 28.628 ARG 1HD 49.685 28.396 ARG 2HD 50.798 27.458 ARG 1HH 49.168 26.032 ARG 1HH1 49.168 26.032 ARG 1HH2 48.173 24.251 ARG 2HH2 47.960 24.316 ASN N 49.996 32.318 ASN CA 50.406 33.706 ASN C 49.508 34.267 ASN O 49.360 33.682 ASN CB 51.893 33.770 ASN CG 52.706 33.250 ASN OD1 52.540 33.681 ASN ND2 53.664 ASN H 50.101 31.681 ASN HA 50.227 34.257 ASN 1HB 52.211 34.789 ASN 2HB 52.099 33.162 ASN 1HD2 54.227 32.167 ASN 2HD2 53.844 32.004 ARG N 48.922 35.407 ARG CA 48.157 36.050 ARG CH 49.030 36.735 ARG CH 49.030 37.689 ARG 2HB 46.686 36.526 ARG 1HG 47.204- 38.380	ARG NE 48.959 26.521 26.267 ARG CZ 48.858 25.909 27.402 ARG NH1 49.281 26.474 28.507 ARG NH2 48.299 24.741 27.436 ARG H 50.337 29.472 21.776 ARG HA 48.004 30.372 23.369 ARG 1HB 49.727 30.401 25.279 ARG 2HB 50.877 29.631 24.213 ARG 1HG 49.197 27.772 23.893 ARG 2HG 48.078 28.628 24.920 ARG 1HD 49.685 28.396 26.850 ARG 2HD 50.798 27.458 25.812 ARG 1HH1 49.168 26.032 29.395 ARG 2HH1 49.705 27.377 28.439 ARG 1HH2 48.173 24.251 28.303 ARG 2HH2 47.960 24.316 26.599 ASN N 49.996 32.318 22.035 ASN CA 50.406 33.706 21.870 ASN C 49.508 34.267 20.783 ASN CB 51.893 33.770 21.499 ASN CB 51.893 33.770 21.499 ASN CB 51.893 33.770 21.499 ASN CB 52.706 33.250 22.622 ASN OD1 52.540 33.681 23.718 ASN CB 51.893 33.770 21.499 ASN CB 51.893 33.770 21.499 ASN CB 51.893 33.770 21.499 ASN CB 52.706 33.250 22.622 ASN OD1 52.540 33.681 23.718 ASN CB 51.893 33.770 21.499 ASN CB 51.893 33.790 ASN CB 51.895 25.407 21.070 ASN CB 51.895 25.407 21.070 ASN CB 51.895 25.407 2	ARG NE 48.959 26.521 26.267 ARG CZ 48.858 25.909 27.402 ARG NH1 49.281 26.474 28.507 ARG NH2 48.299 24.741 27.436 ARG H 50.337 29.472 21.776 ARG HA 48.004 30.372 23.369 ARG 1HB 49.727 30.401 25.279 ARG 2HB 50.877 29.631 24.213 ARG 1HG 49.197 27.772 23.893 ARG 2HG 48.078 28.628 24.920 ARG 1HD 49.685 28.396 26.850 ARG 2HD 50.798 27.458 25.812 ARG 1HH1 49.168 26.032 29.395 ARG 2HH1 49.705 27.377 28.439 ARG 2HH2 48.795 27.377 28.439 ARG 2HH2 47.960 24.316 26.599 ASN N 49.996 32.318 22.035 ASN CA 50.406 33.706 21.870 ASN C 49.508 34.267 20.783 ASN C 49.508 34.267 20.783 ASN C 49.508 34.267 20.783 ASN C 51.893 33.770 21.499 ASN C 65.2706 33.250 22.622 ASN ND 53.664 32.406 22.420 ASN H 50.101 31.681 21.265 ASN HA 50.227 34.257 22.794 ASN 1HB 52.211 34.789 21.274 ASN 2HB 52.099 33.162 20.620 ASN 1HD 54.227 32.167 23.194 ASN 2HB 52.099 33.162 20.620 ASN 1HB 52.211 34.789 21.274 ASN 2HB 52.099 33.162 20.620 ASN 1HB 52.211 34.789 21.274 ASN 2HB 52.099 33.162 20.620 ASN 1HD2 54.227 32.167 23.194 ASN 2HB 52.099 33.162 20.620 ASN 1HD2 54.227 32.167 23.194 ASN 2HB 52.091 33.681 17.785 ARG C 49.030 36.735 18.938 ARG C 49.500 37.866 19.869 ARG CD 45.810 38.876 20.590 ARG NE 44.644 38.387 21.426 ARG CH 43.385 38.416 19.382 ARG NH 43.385 38.416 19.382 ARG NH 43.385 38.316 19.382 ARG NH 43.385 38.316 19.382 ARG NH 48.857 35.623 22.040 ARG HA 47.516 35.294 19.567 ARG CH 47.504 38.380 19.156

					400	
					438	
719	ARG	1HH1	42.596	37.960	18.878	20.00
720	ARG	2HH1		38.855	18.925	20.00
721	ARG		41.874	36.942	20.748	20.00
722	ARG	2HH2	42.724	37.024	22.270	20.00
723	TYR	Ν	50.170	37.367	19.339	12.06
724	TYR	CA	51.144	38.065	18.550	10.98
725	TYR	C	52.522	37.481	18.816	13.59
726	TYR	0	52.966	37.428	19.960	15.53
727	TYR	CB	51.220	39.576	18.875	9.94
728	TYR	CG	49.859	40.250	18.678	11.54
729	TYR	CD1	49.044	40.145	17.523	11.75
730	TYR	CD2	49.380	40.989	19.752	10.70
731	TYR.	CE1	47.808	40.784	17.473	12.58
732	·TYR	CE2	48.142	41.590	19.752	11.42
733	TYR	CZ	47.339	41.494	18.578	12.61
734	TYR	ОН	46.051	41.986	18.423	13.20
735	TYR	Н	50.266	37.312	20.329	20.00
736	TYR	HA	50.862	37.907	17.513	20.00
737	TYR	1HB	51.984	40.094	18.317	20.00
738	TYR	2HB	51.550	39.711	19.903	20.00
739	TYR	HD1	49.336	39.506	16.724	. 20.00
740	TYR	HD2	49.993	41.106	20.635	20.00
741	TYR	HE1	47.172	40.704	16.611	20.00
742	TYR	HE2	48.011	42.008	20.763	20.00
743	TYR	НН	45.857	42.735	18.985	20.00
744	ARG	N	53.153	37.103	17.699	13.40
745	ARG	CA	54.543	36.623	17.580	16.28
746	ARG	C	55.474	37.666	18.247	14.00
747	ARG	0	56.454	37.285	18.895	16.44
748	ARG	CB	54.915	36.428	16.068	17.48
749	ARG	CG	56.305	36.675	15.419	23.38 28.56
750 751	ARG	CD NE	56.316 55.326	37.199 36.527	13.926 13.052	32.94
751 752	ARG	CZ	54.135	36.980	12.495	34.20
753 ·	ARG	NH1	53.971	38.258	12.223	33.15
754	ARG	NH2	53.182	36.079	12.237	33.58
755	ARG	H	52.559	37.155	16.908	20.00
756	ARG	HA	54.579	35.718	18.147	20.00
757	ARG	1HB	54.506	35.533	15.621	20.00
758	ARG	2HB	54.611	37.441	15.734	20.00
759 .	ARG	1HG	56.835	37.432	16.015	20.00
760	ARG	2HG	56.937	35.799	15.540	20.00
761	ARG	1HD	56.210	38.289	13.888	20.00
762	ARG	2HD	57.315	36.970	13.512	20.00
763	ARG	HE -	55.560	35.563	12.998	20.00
764	ARG	1HH2	52.331	36.288	11.752	20.00
765	ARG	2HH2	53.301	35.150	. 12.559	20.00
766	ARG	1HH1	53.238	38.710	11.709	20.00
	-ARG		54.784	- 38.793	12.502	20.00
768	ASP	N .	55.116	38.934	18.094	12.94
769	ASP	CA.	55.973	40.036	18.616	12.52
770	ASP	С	55.797	40.445	20.096	12.63

•			, .			,
771	ASP	0	56.452	41.331	20.648	11.75
772	ASP	СВ	55.837	41.278	17.706	13.44
773	ASP	CG	56.220	41.009	16.226	15.48
774	ASP	OD1	57.040	40.145	15.911	14.91
		OD2		41.579	15.347	17.68
775	ASP		55.608			
776	ASP	Н	54.485	39.225	17.376	20.00
777	ASP	HA	56.988	39.663	18.528	20.00
778	ASP	1HB	56.501	42.075	18.023	20.00
779	ASP	2HB	54.831	41.662	17.752	20.00
780	VAL	N	54.876	39.733	20.803	12.12
781	VAL	CA	54.573	40.057	22.191	11.29
782	VAL	С	54.455	38.762	22.960	9.78
783	VAL	Ο.	53.550	37.945	22.857	11.48
784	VAL	CB	53.595	41.270	22.522	16.11
785	VAL	CG1	52.686	41.008	23.691	15.44 .
786	VAL	CG2	53.024	42.155	21.436	14.33
787	VAL	Н	54.326	39.100	20.257	20.00
788	VAL.	HA	55.521	40.473	22.533	20.00
789	VAL	НВ	54.260	42.006	22.986	20.00
790	VAL	1HG1	53.241	40.703	24.580	20.00
791	VAL		51.975	40.213	23.479	20.00
792	VAL		52.106	41.885	23.978	20.00
793	VAL		52.256	41.649	20.862	20.00
794	VAL		53.807	42.481	20.755	20.00
795	VAL		52.569	43.058	21.839	20.00
796	SER	N	55.506	38.663	23.758	10.16
797	SER	CA	55.834	37.561	24.654	11.16
798	SER	C	56.196	38.081	26.070	9.64
799	SER	0 .	56.758	39.170	26.254	11.51
800	SER	CB	57.105	36.786	24.050	10.81
801	SER	OG	56.905	36.279	22.695	12.18
802	SER	Н	56.131	39.440	23.769	20.00
803	SER	HA	54.941	36.921	24.712	20.00
804	SER	1HB	57.272	35.928	24.740	20.00
						20.00
805	SER	2HB	58.081	37.304	24.285	
806	SER	HG	56.412	36.740	21.950	20.00
807	PRO	N	55.880	37.202	27.077	8.67
808	PRO	CA	56.329	37.300	28.414	10.79
809	PRO	C	57:824	36.995	28.517	13.35
810	PRO	0	58.237	35.943	28.085	14.81
811	PRO	CB	55.432	36.259	29.103	10.00
812	PRO	CG	55.263	35.208	28.112	10.77
813	PRO	CD	55.042	36.035	26.923	9.16
814	PRO	HA	56.203	38.308	28.764	20.00
815	PRO	1HB	54.379	36.441	29.197	20.00
816	PRO	2HB	55.825	35.933	30.055	20.00
817	PRO	1HG	56.174	34.610	28.028	20.00
818	PRO	2HG	54.446	34.516	28.338	20.00
819	-PRO-	1HD-	54.004	36.311	26.766	20.00
820	PRO	2HD	55.499	35.527	26.092	20.00
821	PHE	N:	58.603	37.926	29.144	13.59
822	PHE	CA	59.917	37.512	29.662	12.71

823 824 825 826 827 828 831 832 833 833 834 835 837 838 839 841 842 843 844 845 851 852 853 854 856 866 867 866 866 866 866 866 866 866 86	P P P P P P P P P P P P P P P P A A A A	COCCCCCHH12HHHHHNCCOCCOOHHH12NCCOCCNCCNHHHHH	59.764 58.819 60.730 60.773 61.171 60.557 61.489 60.848 61.354 58.157 60.489 61.767 60.716 61.347 60.194 61.915 60.707 60.799 60.799 60.796 60.121 62.097 60.970 63.206 61.362 59.915 62.233 63.005 61.645 61.906 60.428 63.111 62.763 62.180 63.026 62.180 63.026 62.180 63.026 62.180 63.026 62.180 63.026 62.180 63.026 62.180 63.026 62.180 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.026 63.026 63.027 63.026 63.026 63.027 63.026 63.026 63.026 63.027 63.026 63.027 63.026 63.027 63.026 63.027 63.026 63.027 63.027 63.027 63.026 63.027	36.228 36.148 38.634 39.907 39.952 41.099 41.196 42.337 42.399 38.782 37.218 38.289 38.738 39.055 41.061 41.252 43.231 43.231 43.231 43.231 43.231 34.068 34.428 33.789 33.232 32.558 32.119 32.476 35.357 33.471 32.436 33.861 35.743 36.197 35.970 36.710 38.950	30.513 31.282 30.385 29.668 28.332 30.342 27.800 29.786 28.543 29.377 28.782 30.303 31.460 27.761 31.361 26.806 30.370 28.141 30.364 31.244 32.811 33.631 30.924 29.567 29.186 28.952 29.624 31.028 31.648 31.009 33.141 34.515 35.245 36.418 34.678 34.481 35.412 33.393 34.920 33.707 32.392 34.956 34.003 35.682	12.31 12.68 9.81 10.06 9.58 9.41 10.26 10.02 8.64 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 11.62 12.07 12.27 15.21 13.40 14.25 15.72 12.79 20.00 20.0
862	HIS	NE2	62.587	40.256	33.707	11.30
864	HIS	HA	62.205	34.800	34.956	20.00
866 867	HIS HIS	2HB HD1	63.521 61.951	36.613 38.686	35.682 36.324	20.00 20.00
868 869	HIS HIS	HD2 HE1	63.513 61.660	38.693 41.067	32.486 35.402	20.00
870	SER	N	59.723	36.834	34.519	13.43
871	SER		-58.552	37.448	35.230	12.13
872	SER	C	57.176	36.863	34.862	12.76
873 874	SER SER	O: CB	56.104 58.525	37.349 38.916	35.336 34.846	13:37 10:16
0/4	SER	CD	J0.J25	30.510	34.040	10.10

875 876 877 878 879 880 881 882 883 884 885 886	SER SER SER SER ARG ARG ARG ARG	OG H HA 1HB 2HB HG N CA C O CB CG	58.653 59.957 58.674 59.441 57.743 57.994 57.244 55.939 55.153 55.744 56.173 56.702	39.016 37.166 37.373 39.363 39.503 38.636 35.805 35.373 34.605 33.997 34.471 33.069	33.381 33.600 36.302 35.292 35.420 32.702 34.022 33.508 34.601 35.513 32.280 32.654	10.34 20.00 20.00 20.00 20.00 20.00 11.78 13.97 13.85 11.86 12.96 13.81
887 888	ARG ARG	CD NE	57.016 57.632	32.181 30.888	31.428 31.874	13.20 14.78
889 890	ARG ARG	CZ NH1	57.094 55.822	29.687 29.488	32.079 31.874	15.00 11.82
891	ARG	NH2	57.838	28.727	32.565	17.64
892	ARG	Н	58.122	35.460	33.703	20.00
893 894	ARG ARG	HA 1HB	55.398 56.888	36.266 34.953	33.193 31.615	20.00 20.00
895	ARG	2HB	55.248	34.372	31.713	20.00
896	ARG	1HG	55.959	32.540	33.249	20.00
897	ARG	2HG	57.586	33.156	33.285	20.00
898	ARG	1HD	57.745	32.661	30.777	20.00
899	ARG	2HD	56.126	31.982	30.828 32.076	20.00 20.00
900	ARG ARG	HE 1HH1	58.614 55.444	30.885 28.561	32.078	20.00
902	ARG	2HH1	55.229	30.225	31.600	20.00
903	ARG	1HH2	57.363	27.856	32.770	20.00
904	ARG	2HH2	58.819	28.824	32.735	20.00
905	ILE	N	53.823	34.541	34.371	14.21
906 907	ILE	CA C	53.015 52.784	33.695 32.276	35.241 34.706	12.58 14.61
908	ILE	Ö	52.704	31.983	33.572	15.26
909	ILE	СВ	51.692	34.404	35.385	13.58
910	ILE	CG1	51.914	35.909	35.670	13.29
911	ILE	CG2	50.687	33.768	36.382	13.64
912	ILE ·	CD1 H	51.819	36.312	37.125 33.611	13.30 20.00
913 914	ILE ILE	п НА	53.450 53.491	35.070 33.658	36.221	20.00
915	ILE	HB	51.217	34.358	34.405	20.00
916	ILE	1HG1	51.085	36.426	35.186	20.00
917	ILE		52.780	36.386	35.222	20.00
918	ILE		51.116	33.664	37.379	20.00
919 920	ILE		50.394 49.777	32.762 34.367	36.074 36.452	20.00 20.00
921	ILE		52.673	35.943	37.695	20.00
922	ILE		50.909	35.966	37.616	20.00
923	ILE		51.834	37.400	37.195	20.00
924	LYS	N	53.090	31.420	35.677 35.485	15.82
925 926	LYS LYS	CA C	52.968 51.581	29.984 29.539	35.485 35.986	15.98 17.60
520		_	J 1.55 I	20.000	55.555	

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927	LYS	0	51.237	29.849	37.111	19.95
928	LYS	СB	54.154	29.264	36.206	17.28
929	LYS	CG	55.459	30.029	36.021	17.24
930	LYS	CD	56.580	29.154	36.468	22.81
931	LYS	CE	57.923	29.820	36.118	26.05
932	LYS	NZ	59.002	28.881	36.461	31.86
933	LYS	· H	53.331	31.776	36.578	20.00
934	LYS	HA	53.041	29.782	34.419	20.00
	LYS	1HB	54.240	28.241	35.827	20.00
935				29.180	37.272	20.00
936	LYS	2HB	53.945			20.00
937	LYS	1HG	55.449	30.949	36.604	
938	LYS	2HG	55.572	30.305	34.972	20.00
939	LYS	1HD	56.527	28.185	35.965	20.00
940	LYS	2HD	56.506	28.962	37.538	20.00
941	LYS	1HE	58.051	30.780	36.631	20.00
942	LYS	2HE	57.975	30.045	35.049	20.00
943	LYS	1HZ	58.861	27.982	35.937	20.00
944	LYS	2HZ	58.998	28.698	37.481	20.00
945	LYS	3HZ	59.917	29.296	36.191	20.00
946	LEU	N	50.885	28.827	35.063	17.25
947	LEU	CA	49.701	28.023	35.305	18.29
948	LEU	С	50.164	26.889	36.218	20.34
949	LEU	Ō	51.156	26.241	35.911	19.70
950	LEU	СВ	49.199	27.455	33.986	18.28
951	LEU	C.G	48.053	28.245	33.290	18.66
952	LEU	CD1	47.989	27.995	31.755	17.13
953	LEU	CD2	47.828	29.708	33.707	14.98
954	LEU	H	51.364	28.759	34.195	20.00
955	LEU	HA	48.963	28.629	35.838	20.00
956	LEU	1HB	48.868	26.443	34.106	20.00
	LEU	2HB	50.029	27.419	33.293	20.00
957		HG	47.157	27.754	33.671	20.00
958	LEU	1HD1		26.933	31.522	20.00
959	LEU		48.064			
960	LEU		48.811	28.517	31.259	20.00
961	LEU		47.039	28.334	31.374	20.00
962	LEU		48.762	30.260	33.691	20.00
963	LEU		47,441	29.754	34.725	20.00
964	LEU		47.112	30.213	33.062	20.00
965	HIS	N	49.467	26.676	37.346	22.86
966	HIS	CA	49.798	25.580	38.220	25.86
967	HIS	C	49.188	24.299	37.604	30.06
968	HIS	0	48.407	23:576	38.179	31.82
969	HIS	CB	49.207	25.838	39.607	26.12
970	HIS	CG	49.625	27.147	40.252	25.12
971	HIS	ND1	48.983	27.623	41.360	25.76
972	HIS	CD2	50.620	28.085	39.869	24.91
973	HIS	CE1	49.574	28.827	41.642	25.52
974	HIS	NE2	50.566	29.138	40.757	26.33
975	HIS	Н	48.586	27.148	37.402	20.00
976	HIS	HA	50.880	25.472	38.282	20.00
977	HIS	1HB	49.461	25.018	40.278	20.00
978	HIS	2HB	48.122	25.837	39.553	20.00
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979	HIS	HD1	48.204		27.201		41.765		20.00
980	HIS	HD2	51.269		27.996		39.010		20.00
981	HIS	HE1	49.292		29.465		42.470		20.00
982	GLN	N	49.608		23.987		36.392		.32.94
983	GLN	CA	49.271		22.663		35.842		36.50
984	GLN	C	50.532		21.955		35.351		37.97
985	GLN-	Õ	51.618		22.541		35.302		37.18
986	GLN	СВ	48.147		22.806		34.854		37.10
987	GLN	CG	48.266		24.086		34.021		40.06
988	GLN	CD	47.360		23.885		32.822		44.06
989	GLN	OE1	47.592	•	22.903		32.119		46.49
990	GLN	NE2	46.323		24.719		32.620		41.49
991	GLN	H	50.243	. '	24.713		35.941		20.00
992	GLN	HA	48.926		21.997		36.641		20.00
993	GLN	1HB	47.226		22.901		35.438		20.00
994	GLN	2HB	48.003		21.889		34.282	•	20.00
995	GLN	1HG	49.283		24.268		33.676		20.00
996	GLN	2HG	47.866		24.901		34.612		20.00
997	GLN	1HE2			24.510		31.912		20.00
998	GLN	2HE2			25.495		33.241	٠.	20.00
999	GLIV	N	50.320		20.649		35.092		41.15
1000	GLU	CA	51.547		19.859		34.850		41.70
1000	GLU	CA	51.834		19.728		33.311		39.31
1001	GLU	0	52.984		19.720		32.860		38.68
1002	GLU	СВ	51.430		18.558		35.655		45.57
1003	GLU	CG	51.679		18.866	. '	37.145		54.22
1005	GLU	CD	50.777		17.977		38.035		60.23
1006	GLU	OE1	49.587	1 :	18.325		38.117		63.36
1007	GLU	OE2	51.253	Ŷ	16.964		38.605		62.48
1008	GLU	H	49.451		20.221		35.343		20.00
1009	GLU	HA	52.415	. 1	20.380		35.254		20.00
1010		1HB	52.138		17.803		35.321		20.00
1011	GLU	2HB	50.436		18.128		35.508		20.00
1012	GLU	1HG	51.411		19.888		37.403	•	20.00
1013	GĽŰ	2HG	52.719		18.720		37.428		20.00
1014	ASP	N	50.713		19.782		32:532		38.53
1015	ASP	CA	50.859		19.700		31.061		37.73
1016	ASP	C .	51.793		20.877		30.583		33.50
	ASP	0	52.946		20.793		30.147		35.46
1018	ASP	СВ	49.374		19.842	•	30.492		43.21
1019	ASP	CG	49.448		20.048	•	28.963		49.96
1020	ASP	OD1	50.358		19.411		28.396		53.55
1021	ASP	OD2	48.668		20.859	•	28.386		52.89
1022	ASP	Н	49.807		19.888		32.924		20.00
1023	ASP	·HA	51.321		18.751		30.761		20.00
1024	ASP	1HB	48.803	-	20.635		30.964		20.00
1025	ASP	2HB	48.830		18.915		30.646		20.00
1026	ASN	Ν.	51.117		22.012		30.774		28.37
-1027 -		CA	51-658-		23.198		30.253		21.99
1028	ASN	С	51.272		24.281		31.204		20.24
1029	ASN	0 -	50.088		24.519		31.310		21.87
1030	ASN	CB	51.062		23.311		28.867		19.04
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1031	ASN	CG	51.901	24.202	27.972	18.04
1032	ASN	OD1	52.747	24.954	28.409	19.25
1033	ASN	ND2	51.670	24.083	26.677	15.06
1034	ASN	H	50.148	21.879	30.992	20.00
	ASN	HA				20.00
1035			52.742	23.147	30.225	
1036	ASN	1HB	50.053	23.709	28.917	20.00
1037	ASN	2HB	50.976	22.339	28.367	20.00
1038	ASN		52.136	24.800	26.133	20.00
1039	ASN		51.105	23.357	26.296	20.00
1040	ASP	N	52.310	24.934	31.781	18.00
1041	ASP	CA	52.359	26.033	32.774	18.78
1042	ASP	С	52.269	27.339	32.022	17.35
1043	ASP	0	52.385	28.381	32.643	18.81
1044	ASP	CB	53.678	26.048	33.673	19.48
1045	ASP	CG	55.010	26.577	33.066	23.10
1046	ASP	OD1	55.075	26.798	31.865	24.35
1047	ASP	OD2	56.022	26.809	33.756	29.58
1048	ASP	Н	53.213	24.652	31.479	20.00
1049	ASP	HA	51.485	25.929	33.416	20.00
1050	ASP	1HB	53.905	25.042	34.027	20.00
1051	ASP	2HB	53.466	26.618	34.583	20.00
1052	TYR	N	52.126	27.268	30.658	16.55
1053	TYR	CA	52.342	28.547	29.854	12.98
1054	TYR	C	51.059	29.382	29.617	12.27
1055	TYR	ŏ	50.072	28.908	29.056	12.81
1056	TYR	СВ	53.128	28.308	28.524	10.36
1057	TYR	CG	53.307	29.633	27.752	12.42
1058	TYR	CD1	54.305	30.529	28.104	10.55
1059	TYR	CD2	52.385	30.073	26.782	11.80
1060	TYR	CE1	54.386	31.783	27.545	11.79
1061	TYR	CE2	52.453	31.353	26.260	10.71
1062	TYR	CZ	53.524	32.198	26.591	9.67
1063	TYR	OH	53.938	33.401	26.004	10.99
1064	TYR	H	52.132	26.366	30.228	20.00
1065	TYR	HA	53.022	29.162	30.449	20.00
1066	TYR	1HB	52.598	27.575	27.914	20.00
1067		2HB	54.096	27.862	28.732	20.00
1067		HD1		30.232	28.857	20.00
	TYR		55.008		26.475	
1069	TYR	HD2	51.588	29.416		20,00
1070	TYR	HE1	55.198	32.442	27.833	20.00
1071	TYR	HE2	51:561	31.484	25.630	20.00
1072	TYR	HH	53.565	33.505	25.119	20.00
1073	ILE	N .	51.111	30.653	30.018	13.06
1074	ILE	CA	50.146	31.725	29.683	12.41
1075	ILE	C	50.996	32.948	29.283	11.32
1076	ILE	0	52.043	33.149	29.862	12.19
1077	ILE	CB .	49.059	32.003	30.788	12.98
1078	ILE	CG1	48.065	33.056	30.196	8.76
1079	ILE		49.639	32.315	32.217	10.91
1080	ILE	CD1	46.906 <sup>-</sup>	33.342	31.116	9.88
1081	ILE	Н	51.869 <sup>-</sup>	30.910	30.612	20.00
1082	ILE	HA	49.636	31.400	28.775	20.00

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1083 1084 1085 1086 1087 1088 1090 1091 1092 1093 1094 1095 1096 1097 1098 1099 1100 1101 1103 1104 1105 1106 1107 1110 1111 1111 1111 1111 1111	ASN ASN ASSN ASSN ASSN ALAA ALA ALA	1HG2 2HG2 3HG2 1HD1 3HD1 N C O C O C O C O C O C O C O C O C O C	48.584 50.271 50.242 48.844 47.243 46.420 46.150 50.614 51.251 50.836 49.889 50.740 51.559 51.733 52.102 49.891 52.326 49.721 50.681 52.677 51.472 51.160 52.418 53.203 50.197 52.104 50.721		31.076 32.732 33.993 33.197 31.487 32.475 33.841 32.423 33.978 33.705 34.994 36.013 36.738 35.386 36.512 37.578 36.212 33.381 34.865 35.743 34.548 36.857 35.294 36.004 37.001 37.231 36.292 36.373 35.256 37.888 35.477	445	30.889 29.240 30.018 32.211 32.577 32.935 32.021 31.431 30.654 28.262 27.951 28.762 26.534 25.985 26.614 24.777 27.666 28.001 26.595 25.844 24.276 24.398 30.161 31.204 31.951 32.229 30.340 30.752		20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 9.16 9.65 10.37 10.07 9.08 9.54 10.96 20.00 20.0
1099 1100 1101 1102 1103	ASN ASN ASN ASN	ND2 H HA 1HB 2HB	52.102 49.891 52.326 49.721 50.681		36.212 33.381 34.865 35.743 34.548		24.777 27.666 28.001 26.595 25.844		9.06 20.00 20.00 20.00 20.00
1105 1106 1107 1108 1109	ASN ALA ALA ALA ALA	2HD2 N CA C	51.947 51.472 51.160 52.418 53.203		35.294 36.004 37.001 37.231 36.292		24.398 30.161 31.204 31.951 32.110		20.00 9.41 10.05 11.58 12.92
1111 1112 1113 1114 1115	ALA ALA ALA ALA	Н	52.104		35.256		30.340		20.00
1116 1117 1118 1119 1120 1121 1122	SER SER SER SER SER SER	CA C O CB OG H	53.663 53.285 52.423 54.393 54.544 51.738		38.902 39.621 40.474 39.958 39.424 39.068		33.299 34.550 34.557 32.386 30.979 32.307		11.23 10.85 12.48 7.55 10.45 20.00
1123 1124 1125 1126 1127 1128 1129	SER SER SER SER LEU LEU	HA 1HB 2HB HG N CA C	54.292 55.364 53.966 53.779 54.089 53.997 54.875		38.048 40.197 41.002 38.963 39.317 40.074 41.328		33.555 32.840 32.539 30.539 35.577 36.831 36.773	•	20.00 20.00 20.00 20.00 11.48 12.53 13.15
1130 1131 1132 1133 1134	LEU LEU LEU LEU	O CB CG CD1 CD2	54.873 56.106 54.509 54.050 53.078 55.183	-	41.326 41.237 39.118 39.236 40.321 39.160		36.773 36.710 37.938 39.386 39.819 40.327		13.65 13.18 15.58 15.45 15.90

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1135	LEU	Н	54.798	38.633	35.444	20.00
1136	LEU	HA	52.955	40.335	37.005	20.00
1137	LEU	1HB	55.596	39.023	37.872	20.00
1138	LEU	2HB	54.181	38.124	37.632	20.00
1139	LEU	HG	53.486	38.316	39.575	20.00
1140	LEU	1HD1	52.149	40.243	39.258	20.00
1141	LEU	2HD1	53.486	41.319	39.654	20.00
	LEU	3HD1	52.812	40.238	40.873	20.00
1143	LEU	1HD2		39.996	40.139	20.00
1144	LEU	2HD2		38.233	40.214	20.00
1145	LEU	3HD2		39.213	41.361	20.00
1146		N-	54.198	42.491	36.870	12.91
1147		CA	54.849	43.764	37.188	13.87
1148		C	54.768	44.035	38.703	16.47
1149	ILE	Ö	53.759	44.202	39.394	16.73
1150	ILE	CB	54.282	44.944	36.382	13.55
1151	ILE	CG1	54.385	44.786	34.837	12.82
1152	ILE	CG2	54.940	46.250	36.798	16.67
1153	ILE	CD1	54.103	43.394	34.279	12.75
1154	ILE	Н	53.200	42.393	36.826	20.00
1155	ILE	HA	55.895	43.661	36.897	20.00
1156	ILE	HB	53.217	45.012	36.617	20.00
1157	ILE	1HG1	55.383	45.077	34.511	20.00
1158	ILE		53.700	45.497	34.362	20.00
1159	ILE		56.026	46.220	36.689	20.00
1160	ILE		54.753	46.467	37.845	20.00
1161	ILE		54.556	47.094	36.229	20.00
1162	ILE	1HD1	53.156	42.977	34.619	20.00
1163	ILE	2HD1	54.904	42.685	34.471	20.00
1164	ILE :	3HD1	54.016	43.465	33.195	20.00
1165	LYS	Ň	55.961	44.052	39.204	16.38
1166	LYS	CA	56.155	44.186	40.594	19.03
	LYS	C	56.848	45.516	40.860	18.79
1168	LYS	Ō	58.065	45.591	40.884	18.39
1169	LYS	СВ	56.939	42.956	40.929	24.89
1170	LYS	CG	56.912	42.866	42.439	36.70
1171	LYS	CD	57.328	41.533	43.051	45.35
1172	LYS	CE	57.186	41.692	44.560	49.89
1173	LYS	NZ	57.538	40.467	45.290	53.29
1174:	LYS	Н	56.745	43.962	38.586	20.00
1175	LYS	HA	55.199	44.202	41.123	20.00
1176	LYS	1HB	57.961	43.013	40.539	20.00
1177	LYS	2HB	56.467	42.Ó81	40.482	20.00
1178	LYS	1HG	55.910	43.036	42.726	20.00
1179	LYS	2HG	57.473	43.692	42.872	20.00
1180	LYS	1HD	58.364	41.333	42.777	20.00
1181	LYS	2HD	56.711	40.725	42.648	20.00
1182	LYS	1HE	56.135	41.919	44.779	20.00
1183		2HE	57.770	42.540	44.932	20.00
1184	LYS	1HZ	58.555	40.282	45.176	20.00
1185		2HZ	57.015	39.665	44.873	20.00
1186	LYS	3HZ	57.314	40.520	46.304	20.00

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1187	MET	Ν	56.027	46.567	41.062	17.44
1188	MET	CA	56.634	47.846	41.393	17.07
1189	MET	С	57.007	47.929	42.923	17.04
1190	MET	Ö	56.147	48.095	43.790	16.87
1191	MET	СВ	55.675	48.907	40.957	15.46
1192	MET	CG	55.291	48.734	39.503	15.47
1193	MET	SD	56.753	48.529	38.450	16.71
1194	MET	CE	57.080	50.195	38.026	12.89
1195	MET	H	55.032	46.464	41.035	20.00
1196	MET	HA	57.546	47.951	40.808	20.00
1197	MET		56.133	49.884	41.087	20.00
		1HB				
1198	MET	2HB	54.772	48.927	41.568	20.00
1199	MET	1HG	54.709	49.588	39.160	20.00
1200	MET	2HG	54.618	47.892	39.385	20.00
1201	MET	1HE	57.389	50.721	38.923	20.00
1202	MET	2HE	56.172	50.653	37.664	20.00
1203	MET	3HE	57.869	50.255	37.283	20.00
1204	GLU	Ν	58.311	47.763	43.214	20.27
1205	GLU	CA	58.778	47.729	44.589	22.54
1206	GLU	С	58.455	49.032	45.367	22.33
1207	GLU	0	57.551	49.145	46.174	24.08
1208	GLU	CB	60.261	47.369	44.550	26.36
1209	GLU	CG	60.752	46.542	45.777	35.25
1210	GLU	CD	62.306	46.400	45.765	40.00
1211	GLU	OE1	63.007	47.377	46.070	43.15
1212	GLU	OE2	62.798	45.322	45.423	43.54
1213	GLU	Н	58.846	47.524	42.399	20.00
1214	GLU	HA	58.235	46.912	45.065	20.00
1215	GLU	1HB	60.894	48.246	44.415	20.00
1216	GLU	2HB	60.457	46.747	43.675	20.00
1217	GLU	1HG	60.308	45.548	45.809	20.00
1218	GLU	2HG	60.494	47.045	46.708	20.00
1219	GLU	N	59.176	50.055	45.053	23.41
1220	GLU	CA	58.964	51.347	45.670	24.81
1221	GLU	C	57.479	51.755	45.865	24.74
1222	GLU	ŏ	57.051	52.142	46.947	24.77
1223	GLU	СВ	59.765	52.321	44.797	26.99
1223	GLU	CG	59.810	53.801	45.255	35.79
1225	GLU	CD	60.222	54.688	44.049	45.06
						48.32
1226	GLU	OE1	59.969	54.335	42.876	
1227	GLU	OE2	60.747	55.769	44.308	49.68
1228	GLU	Н	59.978	49.853	44.499	20.00
1229	GLU	HA	59.422	51.306	46.657	20.00
1230	GLU	1HB	59.378	52.294	43.798	20.00
1231	GLU	2HB	60.779	51.944	44.681	20.00
1232	GLU	1HG	60.583	53.892	46.009	20.00
1233	GLU	2HG	58.858	54.133	45.660	20.00
1234	ALA	N	56.670	51.652	44.799	22.17
1235	ALA	CA	55.246	52.057	44.836	21.64
1236	ALA	С	54.359	51.006	45.549	22.46
1237		0	53.160	51.123	45.691	24.05
1238	ALA	СВ	54.806	52.202	43.399	20.53

						448	3		
1239	ALA	Н	57.053	5	51.248		43.967		20.00
1240	ALA	HA	55.174		3.008		45.366		20.00
1241	ALA	1HB	54.756		51.228		42.929		20.00
1242	ALA	2HB	55.522		2.775		42.815		20.00
1243	ALA	3HB	53.829	_	2.664		43.317		20.00
1244	GLN	N	55.030		9.944		45.995		23.74
1245	GLN	CA	54.391		18.860		46.644		27.80
1246	GLN	C	53.125		18.391		45.853		26.92
1247	GLN	Ö	52.149		7.989		46.495		27.71
1247	GLN	СВ	54.140		9.267		48.138		32.88
1249	GLN	. CG	55.326		9.006		49.110		40.36
1250	GLN	CD	55.212		7.532		49.760		47.19
1251	GLN	OE1			7.292		50.673		52.84
			54.421						
1252	GLN	NE2	56.012		6.546		49.265		46.59
1253	GLN	H	55.964		9.826		45.665		20.00
1254	GLN	HA	55.085		8.022		46.580		20.00
1255	GLN	1HB	53.282		8.717		48.513		20.00
1256	GLN	2HB	53.816		0.302		48.171		20.00
1257	GLN	1HG	55.368		9.735		49.918		20.00
1258	GLN	2HG	56.264		9.080		48.553		20.00
1259	GLN	1HE2	55.903		5.690		49.757		20.00
1260	GLN	2HE2			6.689		48.503		20.00
1261	ARG	N	53.166		8.361		44.464		23.43
1262	ARG	CA	52.057		7.713		43.751	;	18.91
1263	ARG	С	52.453		6.726		42.694		16.84
1264	ARG	0_	53.318		7.057		41.916		17.96
1265	ARG	CB	51.320		8.807		43.076		18.58
1266	ARG	CG	50.098		8.257		42.395		16.36
1267	ARG	CD	49.034		9.322	1.25	42.305		17.70
1268	ARG	NE	48.208		9.346		43.527		17.29
1269	ARG	CZ	47.441		0.424		43.806		15.94
1270	ARG	NH1	47.297		1.403		42.891		13.68
1271	ARG	NH2	46.819		0.417		44.975		18.40
1272	ARG	Н	54.018		8.616		44.002		20.00
1273	ARG	HA	51.427		7.185		44.462		20.00
1274	ARG	1HB	51.976		9.317		42.372		20.00
1275		2HB	51.053		9.558		43.825		20.00
1276	ARG	1HG	49.683		7.403		42.929		20.00
1277	ARG	2HG	50.353		7.899		41.397		20.00
1278	ARG	1HD	48.372		9.152		41.452		20.00
1279	ARG	2HD	49.446		0.319		42.247		20.00
1280	ARG	HE	48.258		8.588	•	44.171		20.00
1281	ARG	1HH1	46.755		2.223		43.004		20.00
1282	ARG	2HH1	47.752		1.289		41.992		20.00
1283	ARG		46.269		1.202		45.245		20.00
1284	ARG		46.898		9.595		45.539		20.00
1285	SER	N	51.766		5.572		42.683		13.82
1286	SER	CA	51.692		4.596		41.633		12.48
1287	SER		50.434		4.551		40.727		11.37
1288	SER	0	49.287		4.615		41.157		13.16
1289	SER	CB	51.817		3.113		42.100		12.86
1290	SER	OG	53.028	4	2.834		42.899		19.01

1291	SER	Н	51.167	45.419	43.464	20:00
1292	SER	HA	52.523	44.871	41.005	20.00
1293	SER	1HB	51.950	42.559	41.136	20.00
1294	SER	2HB	50.804	42.672	42.377	20.00
1295	SER	HG	53.168	43.269	43.803	20.00
1296	TYR	N	50.787	44.235	39.430	11.99
1297	TYR	CA	49.732	44.002	38.418	10.12
1298	TYR	C	50.149	42.759	37.675	10.12
1299	TYR	Ö	51.329	42.739	37.602	11.32
1300	TYR	СВ	49.562	45.151	37.002	10.39
				46.542		
1301	TYR	CG	49.764		37.950	10.08
1302	TYR	CD1	51.063	46.952	38.255	10.55
1303	TYR	CD2	48.695	47.416	38.159	10.53
1304	TYR	CE1	51.305	48.211	38.707	12.90
1305	TYR	CE2	48.895	48.689	38.649	12.10
1306	TYR	CZ	50.180	49.106	38.885	13.13
1307	TYR	ОН	50.199	50.433	39.253	13.22
1308	TYR	Н	51.751	44.131	39.179	20.00
1309	TYR	HA	48.782	43.799	38.924	20.00
1310	TYR	1HB	48.582	45.095	36.888	20.00
1311	TYR	2HB	50.285	45.040	36.555	20.00
1312	TYR	HD1	51.873	46.248	38.135	20.00
1313	TYR	HD2	47.680	47.117	37.910	20.00
1314	TYR	HE1	52.383	48.243	38.882	20.00
1315	TYR	HE2	48.058	49.352	38.782	20.00
1316	TYR	НН	50.979	50.728	39.691	20.00
1317	ILE	N	49.222	42.018	37.068	10.35
1318	ILE	CA	49.503	41.053	35.972	7.97
1319	ILE	C	49.087	41.676	34.586	9.78
1320	ILE	Ö	47.914	41.999	34.387	10.25
1321	ILE	СB	48.760	39.736	36.289	9.05
1322		CG1	49.269	39.172	37.681	10.20
1323	ILE	CG2	48.976	38.754	35.095	8.37
1324	ILE	CD1	48.737	37.834	38.236	8.33
1325		H	48.275	42.270	37.213	20.00
1326		HA	50.572	40.865	35.958	20.00
1327		HB.	47.694	39.953	36.380	20.00
1328	ILE		49.048	39.919	38.429	20.00
1329	ILE		50.359	39.121	37.645	20.00
1330	ILE		50.043	38.615	34.936	20.00
		•	48.543			20.00
1331	ILE			39.117	34.167	
1332	ILE		48.537	37.783	35.302	20.00
1333	ILE		49.021	37.050	37.542	20.00
1334	ILE		47.661	37.804	38.225	20.00
1335	ILE		49.128	37.549	39.212	20.00
1336	LEU	N	50.086	41.866	33.670	8.73
1337	LEU	CA	49.787	42.333	32.269	8.66
1338	LEU	С	49.661	41.053	31.404	11.10
1339		0	50.565	40.223	31.418	12.49
	LEU	CB	50.852	43.343	31.645	9.65
1341	LEU	CG	50.557	44.804	32.004	10.58
1342	LEU	CD1	51.628	45.857	31.638	11.19

1343 1344 1345 1346 1347 1348 1350 1351 1352 1353 1354 1355 1356 1357 1358 1360 1361 1362 1363 1364 1365 1366 1367 1376 1377 1378 1379 1370 1371 1372 1373 1374 1375 1376 1377 1378 1379 1380 1381 1382 1383 1384 1385 1386 1387	UUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUU	3HD2 NCCOCCHHHH1123NCCOCCONHABGGG22 NCCOCCHABGGG2 NCCOCCONHABGGG22 NCCOCCONHABGGG22 NCCOCCONHABGGG22 NCCOCCONHABGGG22	49.440 50.173 48.531 48.266 47.699 47.222 47.333 47.456 45.834 47.838 49.265 47.726 47.258 45.238 45.766 45.335 47.760 47.071 45.534 45.108 47.562 47.407 47.989 47.581 49.116 48.093 47.281 48.615 47.022 46.374 47.973 49.523 49.492 44.700 43.381	44.924 41.592 42.831 43.235 43.080 45.070 45.864 46.862 44.547 44.385 45.958 40.850 39.641 40.039 41.124 38.650 37.292 39.019 41.569 39.242 38.689 39.174 39.973 38.315 39.610 39.436 39	33.515 33.979 32.284 30.561 32.000 31.499 30.559 32.073 31.958 34.051 33.809 30.708 29.888 28.466 28.225 30.660 30.225 30.813 30.829 29.728 31.711 29.323 29.907 31.337 31.484 27.461 26.199 26.344 27.188 25.030 25.399 24.452 23.918 27.639 24.452 23.918 27.639 24.452 23.918 27.639 24.452 23.918 25.596 26.297 23.695 25.468 25.333		12.47 20.00
1388 · 1389 1390	GLY GLY GLY	С О Н	43.447 44.117 45.110	37.969 37.488 40.695	25.067 24.141 24.840		10.89 12.56
1390 1391		H 1HA	45.110 42.923	40.695 40.000	 24.840 24.471		20.00
1392	GLY	2HA	42.825	39.769	26.226	•	20.00
1393 1394	PRO PRO	N CA	42.703 42.702	37.161	25.868		11.90 <sup>-</sup>
1394	rku	CA	42.702	35.712	25.665		10.45

1395 1396 1397 1398 1399 1400 1401 1403 1404 1405 1406 1407 1408 1409 1410 1411 1412 1423 1424 1425 1426 1437 1438 1436 1437 1438 1439 1440 1441 1442 1443 1444 1443		3HD2 N CA C O CB CCD HAB 2HB 1HD 2HD N CA C O	43.657 45.373 40.810 39.625 39.955 39.063 39.185 39.775 41.112 38.865 38.100 39.579 39.869 39.170 41.526 41.807 41.232 41.785 42.556 43.108	35.222 35.804 35.253 36.340 37.636 35.498 34.304 35.140 36.362 36.169 38.326 38.092 34.139 32.441 32.179 32.668 33.411 32.479 32.668 33.411 34.682 33.945 34.195 32.178 31.757 32.734 32.381 34.045 33.682 35.528 34.594 34.903 31.845 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.784 31.757 32.881 31.757 32.882 33.682 31.705 31.845 31.705 31.845 31.705 31.845 31.705 31.845 31	24.206 23.503 26.615 27.662 26.938 26.026 27.113 26.128 28.234 28.380 27.629 26.560 23.738 22.471 22.773 23.932 21.783 21.125 21.981 20.356 24.174 21.784 20.906 22.250 20.321 22.362 22.844 21.388 21.034 19.801 19.634 21.721 21.951 22.707 23.267 20.513 19.619 20.286 22.479 20.422 20.211 18.588 19.648 20.005 19.988 22.729 23.484 24.770 25.417	12.59 12.30 10.51 11.89 9.07 20.00 20.00 20.00 20.00 20.00 20.00 13.19 14.02 13.93 12.95 13.28 12.21 20.00 2
1442	ASN	С	42.556	28.687	24.770	15.36
1444	ASN	CB	42.877	27.633 27.529	23.417	14.48
1445	ASN	CG	44.147	28.314	22.038	16.03
1446	ASN	OD1	44.043	29.255	21.263	19.68
				-		

1447 1448	ASN ASN	ND2 H	45.329 41.818		27.892 29.895		22.449 22.060		15,67 20.00
1449	ASN	HA	40.954		27.643		23.757		20.00
1450 1451	ASN ASN	1HB 2HB	42.375 43.249		27.105 26.672		21.737 23.187		20.00 20.00
1452	ASN	1HD2			28.521		22.054		20.00
1453	ASN	2HD2			27.139		23.052		20.00
1454	THR	Ν	42.650		29.978		25.133		15.02
1455	THR	CA	43.391		30.291		26.417		12.29
1456	THR	C	42.474		31.036		27.517		11.96
1457 1458	THR THR	O CB	42.852	•	31.476		28.583		11.69
1459	THR	OG1	44.719 44.565	4.	31.008 32.343		26.014 25.555		10.87 10.92
1460	THR	CG2	45.584		30.273		25.027	•	10.32
1461	THR	Н	42.262		30.664		24.515		20.00
1462	THR	HA	43.646		29.369		26.917		20.00
1463	THR	НВ	45.361		30.992		26.941		20.00
1464	THR	HG1	43.754		32.671		25.107		20.00
1465 1466	THR THR		45.153 45.915		30.085 29.312		24.039 25.408		20.00
1467	THR		46.499		30.848		24.857		20.00 20.00
1468	CYS	N	41.161		31.075		27.244		14.04
1469	CYS	CA	40.171		31.667		28.138		14.17
1470	CYS	С	40.085		30.933		29.493		12.97
1471	CYS	0	39.963		31.492		30.591		13.90
1472	CYS	CB	38.786		31.617		27.536	,	13.16
1473 1474	CYS	SG H	38.550		32.838		26.281		15.24
1474	CYS CYS	П НА	40.878 40.460		30.770 32.696		26.336 28.339		20.00 20.00
1476	CYS	1HB	38.031		31.820	·	28.305		20.00
1477	CYS	2HB	38.541		30.623		27.152		20.00
1478	CYS	HG	38.695		32.291		25,082		20.00
1479	GLY	N	40.243		29.645		29.298		13.25
1480	GLY	CA	40.387		28.785		30.429		14.11
1481 1482	GLY GLY	C	41.808		28.832		31.180	•	14.77
1483	GLY	Н	41.877 40.296	•	28.613 29.275		32.340 28.376		15.71 20.00
1484	GLY	1HA	40.315		27.866		29.861		20.00
1485	GLY	2HA	39.548		29.026		31.088		20.00
1486	HIS	Ν	42.939		29.144°	•	30.573		15.04
1487	HIS	CA	44.200		29.473	,	31.244		,13.49
1488	HIS	C .	43.971		30.831		31.942		14.38
1489 1490	HIS HIS	O CB	44.520 45.303		31.190 29.659		32.974		15.19
1491	HIS	CG	45.443		28.449		30.149 29.222		12.89 14.63
1492	HIS	ND1	45.683		28.487		27.875		13.48
1493	HIS	CD2	45.237		27.096		29.517		16.55
1494	HIS	CE1	45.572		27.255		27.426		14.33
1495		NE2	45.317		26.382		28.410		15.87
1496	HIS	H	42.914		28.968		29.590		20.00
1497 1498	HIS HIS	HA 1HB	44.420 46.257		28.687 29.992		31.970 30.558		20.00 20.00
	•		1		_0.002		55.550		_0.00

1499 1500 1501 1502 1503 1504 1505 1506 1507 1508 1509 1510 1511 1513 1514 1515 1521 1522 1523 1524 1525 1526 1527 1529 1530 1531 1532 1533 1533 1533 1533 1543 1543 1543 1544 1543 1544	PHE	2HHHN CCOCCCCCHH112HHHHHN CCOCCCCCCCCHH112HHHHN CCOCCCCCCHH114HD1132	44.992 45.880 44.997 45.646 43.159 43.089 42.415 42.872 42.315 42.032 40.875 42.919 40.653 42.743 44.103 41.535 42.743 44.103 41.355 42.841 40.158 43.281 41.317 41.278 40.450 41.113 41.043 39.015 38.226 37.724 37.952 37.180 37.724 37.952 37.180 37.724 37.952 37.180 37.724 37.952 37.724 37.952 37.180 37.724 37.952 37.7265 40.399 38.472 39.104 37.760 36.707 38.784 36.450	30.483 29.274 26.700 26.975 31.695 33.072 33.006 33.545 33.909 35.335 35.627 36.384 36.859 37.665 37.845 31.410 33.467 33.453 33.912 34.849 36.183 37.060 38.521 38.835 32.289 32.222 31.433 31.813 31.707 32.858 32.964 34.113 34.195 34.931 34.589 36.217 35.900 36.217 35.900 36.217 35.900 36.217 35.900 36.217 35.900 36.217 35.900 36.217 36	29.501 27.336 30.496 26.381 31.311 31.837 33.201 34.203 30.774 31.183 30.909 32.368 31.367 32.118 30.457 31.947 30.563 29.814 32.083 30.347 32.953 31.136 32.490 33.173 34.411 35.595 36.756 34.051 31.900 33.399 32.065 34.051 31.900 33.399 32.065 34.051 31.900 33.399 32.065 34.051 31.900 33.399 32.065 34.051 31.900 33.399 32.065 34.051 31.900 33.415 35.617 34.699 32.302 34.783 34.862 33.295 31.345 31.085 36.026 32.687	20.00 20.00 20.00 12.72 12.68 13.22 12.12 12.82 9.60 8.18 8.54 11.79 11.03 10.70 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 12.17 12.15 12.77 12.89 13.28 13.46 13.99 12.69 13.76 11.54 11.61 20.00 20.
1541	TRP	HD1	37.760	32.155	31.345	20.00
1542	TRP	HE1	36.707	34.474	31.085	20.00
1544	TRP	HZ2	36.450	36.835	32.687	20.00
1545	TRP	HZ3	38:094	36.300	36.599	20.00
1546	TRP	HH2	36.986	37.699	34.976	20.00
1547	GLU	N	41.780	30.326	35.178	13.48
1548	GLU	CA	42.730	29.627	35.996	12.78
1549	GLU	C	43.768	30.566	36.534	13.54
1550	GLU	O	43.972	30.594	37.732	13.56

1590 1591 1592 1593 1594 1595 1596 1597 1598	GGGGGGGGGGMMMMMMMMMMMMMMMWWWVVVVVVVVVVV	2HG1 3HG1 1HG2 2HG2 3HG2 N CA C O	43.388 44.436 45.202 44.936 46.076 41.624 42.211 43.840 42.624 43.929 45.161 44.363 44.716 45.324 46.127 46.844 47.623 46.176 46.154 45.456 46.59 47.575 46.59 47.575 46.59 47.575 46.520 43.734 41.931 40.524 43.927 42.614 43.927 42.614 43.927 42.614 43.927 43.927 43.927 43.927 44.0927 42.614 43.927 43.937	28.531 27.771 26.754 26.639 26.069 30.031 29.204 28.925 27.820 27.252 28.434 31.442 32.380 33.407 33.773 33.111 34.402 35.297 36.105 31.230 31.771 33.358 32.430 34.147 35.105 36.709 35.347 36.723 33.877 36.723 33.877 36.723 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.822 37.8222 37.822	35.219 36.130 35.308 34.112 35.793 34.237 36.855 34.313 34.904 36.939 36.597 35.736 36.339 37.35.218 35.674 34.348 33.568 34.763 36.918 34.395 34.785 36.441 36.154 36.395 37.215 36.095 37.215 36.095 37.215 36.095 37.215 36.095 37.215 36.109 38.158 37.215 36.109 38.158 37.215 36.109 38.158 37.215 36.109 38.158 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 37.215 38.356 38.356 37.215 38.356	11.88 15.11 17.43 19.12 19.81 20.00
1598	TRP	0	43.328	32.070	42.288	16.22
1600	TRP	CG	40.867	29.820	41.069	17.46
1601 1602	TRP TRP	CD1 CD2	41.606 39.602	28.897 29.940	41.786 41.731	21.61 19.79
1002	LIM		JJ.UUZ	20.040	71.731	10.15

		,				
1603	TRP	NE1	40.899	28.456	42.861	23.85
1604	TRP	CE2	39.653	29.065	42.870	22.70
1605	TRP	CE3	38.464	30.605	41.428	21.09
1606	TRP	CZ2	38.576	28.895	43.700	21.76
1607	TRP	CZ3	37.360	30.412	42.267	21.15
		CH2	37.413	29.570	43.399	21.23
1608	TRP				38.246	20.00
1609	TRP	Н	42.074	32.302		
1610	TRP	HA	41.294	32.332	40.966	20.00
1611	TRP	1HB	42.030	29.916	39.289	20.00
1612	TRP	2HB	40.487	30.754	39.199	20.00
1613	TRP	HD1	42.618	28.612	41.516	20.00
1614	TRP	HE1	41.220	27.834	43.554	20.00
1615	TRP	HE3	38.407	31.218	40.555	20.00
1616	TRP	HZ2	38.667	28.250	44.569	20.00
1617	TRP	HZ3	36.437	30.916	42.045	20.00
1618	TRP	HH2	36.555	29.455	44.032	20.00
1619	GLU	N	44.259	30.940	40.543	16.06
1620	GLU	CA	45.592	30.695	41.135	15.05
1621	GLU	C	46.223	31.929	41.826	16.50
1622	GLU	Ö	46.831	31.834	42.894	18.37
1623	GLU	СВ	46.473	30.020	40.099	13.43
1624	GLU	CG	45.967	28.568	39.973	13.77
1625	GLU	CD	46.558	27.837	38.747	14.43
1626	GLU	OE1	47.458	28.382	38.119	15.87
	GLU	OE2	46.113	26.745	38.435	15.61
1627				30.564	39.620	20.00
1628	GLU	Н	44.114	29.979	41.932	20.00
1629	GLU	HA	45.422		40.407	20.00
1630	GLU	1HB	47.522	30.030		20.00
1631	GLU	2HB	46.364	30.567	39.152	
1632	GLÜ	1HG	44.895	28.546	39.840	20.00
1633	GLU	2HG	46.191	27.991	40.858	20.00
1634	GLN	N	45.999	33.062	41.202	14.99
1635	GLN	CĄ	46.714	34.258	41.610	14.10
1636	GLN .	С	45.841	35.103	42.496	14.79
1637	GLN	0	46.292	36.113	42.957	14.03
1638	GLN	CB	47.074	35.107	40.341	13.84
1639	GLN	CG	48.001	34.423	39.298	14.77
1640	GLN	CD	49.291	33.960	40.073	20.12
1641	GLN	OE1	49.843	34.780	40.814	21.85
1642	GLN	NE2	49.740	32.664	39.964	18.11
1643	GLN	Н	45.582	33.013	40.294	20.00
1644	GLN	HA	47.620	34.012	42.178	20.00
1645	GLN	1HB	47.560	36.034	40.653	20.00
1646	GLN	2HB	46.157	35.450	39.856	20.00
1647	GLN	1HG	48.269	35.110	38.493	20.00
1648	GLN	2HG	47.536	33.553	38.850	20.00
1649	GLN		50.530	32.405	40.514	20.00
1650	GLN		49.411	31.946	39.364	20.00
	- LYS-		-44.583 ···	- 34.682	42.756	14.24
1652	LYS	CA	43.720	35.355	43.752	15.01
			43.720	36.821	43.456	12.96
1653	LYS	C		37.665	43.430	13.31
1654	LYS	0	43.245	37.003	. 44,331	13.51

1655	LYS	СВ	44.218	35.116	45.226	19.21
1656	LYS	CG	44.427	33.587	45.521	21.85
1657	LYS	CD	44.580	33.241	47.008	29.47
1658	LYS	CE	45.043	31.802	47.295	31.82
1659	LYS	NZ	46.358	31.625	46.658	37.86
1660	LYS	Н	44.246	33.917	42.208	20.00
1661	LYS	HA	42.775	34.834	43.635	20.00
1662	LYS	1HB	43.461	35.509	45.903 45.430	20.00
1663-	LYS	2HB	45.135	35.675 33.231	45.430 44.929	20.00
1664.	LYS LYS	1HG 2HG	45.269 43.538	33.231	45.155	20.00
1665 1666	LYS	1HD	43.644	33.429	47.532	20.00
1667	LYS	2HD	45.293	33.928	47.452	20.00
1668	LYS	1HE	44.353	31.026	46.952	20.00
1669	LYS	2HE	45.156	31.662	48.378	20.00
1670	LYS	1HZ	47.041	32.309	47.036	20.00
1671	LYS	2HZ	46.295	31.809	45.636	20.00
1672	LYS	3HZ	46.730	30.665	46.798	20.00
1673	SER	N	43.292	37.085	42.172	14.72
1674	SER	CA	42.944	38.424	41.776	14.22
1675	SER	С	41.469	38.644	41.990	15.12
1676	SER	0	40.653	37.731	41.873	14.48
1677	SER	CB	43.222	38.649	40.258	12.78
1678	SER	OG	44.599	38.384	39.786	13.54
1679	SER	Н	43.501	36.368	41.511	20.00
1680	SER	HA	43.424	39.132	42.463	20.00
1681	SER	1HB	42.684	39.566	39.862	20.00
1682	SER	2HB	42.553	37.916	39.760	20.00
1683	SER	HG	45.445	38.774	40.181	20.00
1684	ARG	N	41.194	39.943	42.269	13.87 14.62
1685	ARG	CA	39.838	40.374 41.125	42.565 41.397	14.02
1686	ARG	C O	39.210 38.037	41.125	41.111	13.72
1687 1688	ARG ARG	CB	39.935	41.251	43.846	14.39
1689	ARG	CG	38.548	41.299	44.523	23.02
1690	ARG	.CD	37.577	42.376	44.071	27.49
1691	ARG	NE	36.501	42.588	45.040	30.20
1692	ARG	CZ	35.270	42.112	45.005	29.90
1693	ARG		34.915	40.949	44.459	31.36
1694	ARG	NH2	34.360	42.901	45.542	30.49
1695	ARG	Н	41.983	40.543	42.411	20.00
1696	ARG	HA	39.228	39.487	42.741	20.00
1697	ARG	1HB	40.353	42.237	43.656	20.00
1698	ARG	2HB	40.624	40.744	44.518	20.00
1699	ARG	1HG	38.665	41.371	45.589	20.00
1700	ARG	2HG	38.050	40.344	44.361	20.00
1701	ARG	1HD	37.164	42.250	43.068	20.00
1702	ARG	2HD	38.127	43.313	44.077	20.00
1703	ARG	HE	36.646	43.361	45.657	20.00
1704	ARG	1HH1	33.929	40.756.	44.381 44.127	20.00 20.00
1705	ARG	2HH1		40.273 42.577	44.127 45.455	20.00
1706	ARG	IMMZ	33.407	42.377	40,400.	20.00

1707	ARG	2HH2	34.560	43.766	45.978	20.00
1708	GLY	N	40.103	41.883	40.761	15.17
1709	GLY	CA	39.769	42.723	39.598	13.10
1710	GLY	С	40.451	42.232	38.273	12.93
1711	GLY	Ö	41.570	41.712	38.267	12.18
1712	GLY	H	41.036	41.839	41.124	20.00
1713	GLY	1HA	40.096	43.738	39.816	20.00
1714	GLY	2HA	38.686	42.743	39.482	20.00
1715	VAL	N.	39.674	42.483	37.168	13.38
1716	VAL	CA	40.227	42.427	35.793	10.87
1717	VAL	C	39.995	43.725	35.172	9.66
1718	VAL	Ö	38.888	44.245	35.106	10.04
1719	VAL	СВ	39.570	41.284	34.982	10.73
1720	VAL	CG1	39.639	39.894	35.612	11.72
1721	VAL	CG2	40.122	41.100	33.537	10.72
1722	VAL	H	38.722	42.761	37.332	20.00
1723	VAL	HA	41.295	42.751	35.853	20.00
1723	VAL	HB.	38.516	41.544	34.929	20.00
1725	VAL	1HG1	39.229	39.859	36.619	20.00
1725	VAL	2HG1		39.558	35.682	20.00
1727	VAL		39.100	39.172	34.999	20.00
1728	VAL		41.197	40.918	33.522	20.00
1729	VAL		39.948	41.984	32.925	20.00
1729	VAL		39.637	40.265	33.032	20.00
1731	VAL	N	41.057	44.256	34.606	8.51
1731	VAL	CA.	40.949	45.502	33.785	7.91
1733	VAL	C	41.098	45.261	32.275	8.39
1734	VAL	Ö	42.171	44.892	31.831	9.71
1735	VAL	СВ	41.905	46.631	34.279	6.14
1736	VAL	CG1	41.684	47.022	35.734	7.70
1737	VAL	CG2	41.778	47.812	33.395	6.32
1738	VAL	H	41.884	43.822	34.895	20.00
1739	VAL	НА	39.938	45.889	33.912	20.00
1740		HB	42.928	46.278	34.202	20.00
1741	VAL		41.809	46.162	36.389	20.00
1742	VAL		40.681	47.425	35.882	20.00
1743	VAL		42.394	47.791	36.054	20.00
1744	VAL		40.751	48.182	33.347	20.00
1745	VAL		42.150°	47.616	32.391	20.00
1746	VAL		42.380	48.634	33.780	20.00
1747	MET	N	40.029	45.551	31.499	8.82
1748	MET	CA	40.098	45.371	30.025	8.53
1749	MET	C	40.050	46.649	29.219	9.29
1750	MET	Ö	39.015	47.259	29.225	9.66
1751	MET	СВ	38.784	44.680	29.671	8.63
1752	MET	CG	38.641	44.332	28.205	9.02
1753	MET	SD	37.443	43.055	27.893	13.59
1754	MET	CE	37.600	42.908	26.065	7.30
1755	MET	H	39.197	45.854	31.962	20.00
1756	MET	HA	40.941	44.743	29.751	20.00
1757	MET	1HB	37.913	45.199	30.067	20.00
1758	MET	2HB	38.900	43.841	30.315	20.00

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1759	MET	1HG	39.586	43.983	27.797	20.00
1760	MET	2HG	38.320	45.209	27.645	20.00
1761	MET	1HE	38.649	42.746	25.840	20.00
1762	MET	2HE	37.313	43.833	25.576	20.00
1763	MET	3HE	37.023	42.066	25.702	20.00
1764	LEU	N	41.108	47.023	28.523	8.05
1765	LEU	CA	41.161	48.369	27.911	6.79
1766	LEU	C	40.878	48.480	26.338	8.29
1767	LEU	Ŏ.	41.027	49.534	25.740	8.17
1768	LEU	СВ	42.531	48.968	28.277	8.82
1769	LEU	CG	42.851	48.939	29.793	9.33
1770	LEU	CD1	41.825	49.803	30.601	6.68
1771	LEU	CD2	44.286	49.352	29.927	8.69
1772	LEU	H	41.914	46.450	28.660	20.00
		HA .				
1773	LEU		40.405	48.977	28.396	20.00
1774	LEU	1HB	42.648	49.982	27.900	20.00
1775	LEU	2HB	43.278	48.364	27.763	20.00
1776	LEU	HG	42.807	47.915	30.157	20.00
1777	LEU	1HD1	40.826	49.357	30.594	20.00
1778		2HD1	41.756	50.793	30.148	20.00
1779	LEU	3HD1	42.127	49.940	31.640	20.00
1780			44.381	50.415	29.708	20.00
1781	LEU		44.933	48.881	29.208	20.00
1782	LEU		44.686	49.148	30.922	20.00
1783	ASN	N	40.414	47.337	25.761	8.85
.1784	ASN	CA	40,131	47.123	24.392	10.85
1785	ASN	C	38.659	46.635	24.231	10.96
1786	ASN	0	38.021	46.324	25.247	11.52
1787	ASN	CB	41.197	46.126	23.941	10.56
1788	ASN	CG	40.890	44.664	24.261	10.18
1789	ASN	OD1	40.432	43.836	23.484	13.30
1790	ASN	ND2	41.203	44.346	25.480	6.84
1791	ASN	Η,	39.979	46.747	26.435	20.00
1792	ASN	ΉA	40.182	48.082	23.880	20.00
1793	ASN	1HB	42.177	46.387	24.344	20.00
1794	ASN	2HB	41.276	46.154	22.858	20.00
1795	ASN	1HD2	41.057	.43.367	25.597	20.00
1796	ASN	2HD2	41.445	44.908	26.261	20.00
1797	ARG	N	38.180	46.553	<b>22.953</b>	12.81
1798	ARG	CA	36.927	45.848	22.626	12.84
1,799	ARG	Č ,	37.350	44:502	22.051	13.06
1800	ARG	0.	38.503	44.343	21.679	13.72
1801	ARG	CB	35.917	46.694	21.830	15.06
1802	ARG	CG ·	35.700	48.081	22.459	23.73
1803	ARG	CD	34.608	48.863	21.755	34.40
1804		NE	34.599	48.708	20.298	44.56
	ARG		35.344	49.461	19.525	51.42
1806	ARG	NH1	36.112	50.366	20.022	54.99
	ARG		35.346	49.337	18.230	53.99
1808	ARG	H	38.768	46.870	22.206	20.00
1809	ARG	HA	36.453	45.621	23.579	20.00
1810	ARG	1HB	34.965	46.180	21.772	20.00
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1811	ARG	2HB	36.260	46.803	20.803	20.00
1812	ARG	1HG	36.626	48.660	22.436	20.00
1813	ARG	2HG	35.439	47.990	23.514	20.00
1814	ARG	1HD	34.562	49.917	22.003	20.00
1815	ARG	2HD	33.643	48.479	22.083	20.00
1816	ARG	HE	34.015	48.023	19.871	20.00
1817	ARG	1HH1	36.646	50.991	19.465	20.00
1818	ARG	2HH1	36.185	50.455	21.013	20.00
1819	ARG	1HH2	35.928	49.944	17.707	20.00
1820	ARG	2HH2	34.791	48.643	17.775	20.00 13.43
1821	VAL	N	36.412	43.522 42.221	22.089 21.443	16.12
1822 1823	VAL VAL	CA C	36.678 36.936	42.221	19.861	16.48
1824	VAL	0	37.508	41.501	19.204	14.56
1825	VAL	СВ	35.416	41.364	21.767	14.79
1826	VAL	CG1	35.507	40.697	23.158	14.80
1827	VAL	CG2	35.222	40.300	20.662	16.58
1828	VAL	Н	35.544	43.637	22.538	20.00
1829	VAL	HA	37.574	41.806	21.888	20.00
1830	VAL	НВ	34.531	42.004	21.742	20.00
1831	VAL	1HG1	35.515	41.454	23.937	20.00
1832	VAL	2HG1	36.431	40.123	23.260	20.00
1833	VAL		34.687	40.009	23.336	20.00
1834	VAL	1HG2	36.123	39.694	20.523	20.00
1835	VAL	2HG2	34.966	40.732	19.690	20.00
1836	VAL	3HG2	34.404	39.618	20.907	20.00
1837	MET	Ν	36.406	43.407	19.280	17.62
1838	MET	CA	36.626	43.711	17.914	19.41
1839	MET	C	36.901	45.186	17.748	17.22
1840	MET	0	36.164	46.011	18.235	16.41
1841	MET	СВ	35.353	43.277	17.172	22.53
1842	MET	CG	35.669	43.239	15.645	30.44
1843	MET	SD	34.456	42.296	14.726	36.43
1844	MET	CE	33.433	43.769	14.332	32.23 20.00
1845	MET	Н	35.921	44.042 43.147	19.872 17.583	20.00
1846 1847	MET MET	HA 1HB	37.494 34.536	43.147	17.353	20.00
1848	MET	2HB	35.027	42.298	17.537	20.00
1849	MET	1HG	36.598	42.694	15.471	20.00
1850	MET	2HG	35.835	44.229	15.216	20.00
1851	MET	1HE	34.049	44.530	13.845	20.00
1852	MET	2HE	33.035	44.226	15.235	20.00
1853	MET	3HE	32.615	43.541	13.649	20.00
1854	GLU	N	37.970	45.511	17.065	17.49
1855	GLU	CA	38.300	46.936	16.903	18.64
1856	GLU	Ċ	38.818	47.158	15.437	17.88
1857	GLU	0	39.395	46.229	14.856	18.32
1858	GLU	СВ	39.271	47.483	17.997	16.54
1859	GLU	- CG	39.163	46.802	19.374	16.82
1860	GLU	CD	40.244	47.335	20.294	15.94
1861	0_0	OE1	41.417	47.016	20.261	15.52
1862	GLU	OE2	39.878	48.107	21.133	15.98

1863 1864 1865 1866 1867 1868 1870 1871 1872 1873 1874 1875 1876 1877 1880 1881 1883 1884 1885 1886 1890 1891 1893 1894 1895 1896 1897 1898 1890 1901 1903 1904 1905 1907 1908 1909 1910 1911	GGGGGGL1111111111111111111111111111111	H H 11 21 2 N C C O C C C N H H 1 21 21 21 21 21 21 N C C O H 1 1 2 N C C O C O H H 1 2 H N C C O H 1 2 H N C C O C O H H 1 2 H N C C O C O H H 1 2 H N C C O C O H H 1 2 H N C C O C O C O C O C O C O C O C O C O	38.603 37.372 39.171 40.290 39.267 38.189 38.446 38.777 38.520 39.311 40.703 42.172 43.021 42.319 37.978 38.115 40.638 40.674 42.152 43.661 41.751 43.043 37.431 37.164 37.364 43.661 41.751 43.09 37.431 37.939 37.431 37.	44.799 47.509 48.563 47.331 45.731 47.010 48.382 48.777 47.522 47.173 49.240 50.654 50.552 51.809 53.111 49.029 49.580 49.186 48.468 50.973 51.644 53.142 53.243 53.887 46.797 45.601 44.317 43.318 47.142 45.858 45.378 44.320 43.056 42.486 43.233 43.073 43.211 45.222 42.275 42.062 43.644 43.991 41.139 40.408	16.761 16.964 18.106 17.643 19.339 19.799 14.874 13.484 12.524 11.670 13.415 13.986 14.578 14.894 14.983 15.463 13.155 12.388 13.917 14.782 13.184 13.889 15.476 14.065 15.777 15.851 14.157 15.036 12.782 11.983 12.298 11.675 13.472 10.936 12.101 13.292 13.660 15.005 15.902 13.410 11.962 13.434 13.020 13.716 14.204 11.368 15.085 16.344	20.00 20.00 20.00 20.00 19.17 20.14 18.74 18.86 21.50 24.11 28.82 30.30 32.58 20.00
1911 1912	LEU -	O 	40.426 41.500	40.584 40.325	17.073 16.507	14.22 14.90 16.30
1913 1914	LEU	CB CG	38.953 37.613	38.890 38.637	16.239 15.495	13.16 12.66

1915 1916 1917 1918 1919 1920 1921 1923 1924 1925 1926 1927 1928 1929 1930 1931 1932 1933 1934 1943 1943 1944 1945 1948 1949 1950 1951 1952 1953 1955 1956 1957 1958 1960 1961 1962 1963 1965	LILUUUUUUUUUUUSSSSSSSSSSSSSSSSSSSSSSSSS	CCH H12H123NCCOCCCNH H121121121121NCCOCSH H12HN COCSH ABB D112112112112112112112112112112112112112	36.136 35.484 37.257 38.130	39.491 37.112 40.665 40.914 38.447 38.405 39.052 40.564 39.341 36.579 36.677 36.951 41.721 40.990 41.101 42.19 44.088 44.879 41.565 43.590 43.359 45.121 43.779 43.070 44.442 45.933 44.495 43.803 45.336 45.200 40.163 39.378 38.345 38.571 40.930 40.163 39.378 38.345 38.571 40.335 39.378 38.345 38.571 40.335 39.378 39.	•	16.079 15.281 14.226 16.899 17.234 15.689 14.496 16.142 17.085 15.469 16.225 14.681 14.742 18.298 19.186 20.578 21.208 19.400 18.204 18.074 16.995 17.227 18.494 18.549 20.273 19.638 17.280 17.823	15.43 13.88 20.00 20.00 20.00 20.00 20.00 20.00 20.00 14.30 11.19 10.51 11.24 11.90 15.22 16.86 20.26 18.45 20.00
1966	ALA	Н	39.961	37.072		23.467	20.00

1967	ALA	НА	37.727	36.169	21.837	20.00
1968	ALA	1HB	38.943	35.142	24.518	20.00
1969	ALA	2HB	39.334	34.524	22.920	20.00
1970	ALA	3HB	37.722	34.257	23.606	20.00
1971	GLN	N	35.725	36.522	23.415	12.25
1972	GLN	CA	34.645	36.830	24.422	13.18
1973	GLN	C	34.789	35.908	25.674	13.71
1974	GLN	Ö	34.306	34.800	25.733	14.92
1975	GLN	СВ	33.424	36.554	23.562	13.64
1976	GLN	CG	32.149	37.021	24.112	14.19
1977	GLN	CD	32.229	38.437	24.551	14.78
1978	GLN	OE1	32.151	38.741	25.725	20.52
1979	GLN	NE2	32.162	39.286	23.542	13.59
1980	GLN	Н	35.602	35.916	22.635	20.00
1981	GLN	HA	34.723	37.882	24.718	20.00
1982	GLN	1HB	33.359	35.487	23.379	20.00
1983	GLN	2HB	33.554	37.017	22.589	20.00
1984	GLN	1HG	31.839	36.429	24.969	20.00
1985	GLN	2HG	31.338	36.929	23.379	20.00
1986	GLN	1HE2		40.195	23.842	20.00
1987	GLN	2HE2		39.058	22.593	20.00
1988	TYR	N	35.597	36.366	26.628	12.19
1989	TYR	CA	36.060	35.379	27.613	11.14
1990	TYR	С	35.168	35.400°	28.882	10.72
1991	TYR	0	35.292	34.588	29.784	12.61
1992	TYR	CB	37.551	35.557	27.877	10.70
1993	TYR	CG	37.814	36.894	28.483	12.19
1994	TYR	CD1	37.736	37.113	29.882	10.21
1995	TYR	CD2	38,194	37.946	27.679	11.16
1996	TYR	CE1	38.060	38.311	30.487	12.37
1997	TYR	CE2	38.525	39.175	28.244	10.83
1998	TYR	CZ	38.452	39.353	29.637	11.09
1999	TYR	ОН	38.741	40.615	30.127	11.70
2000	TYR	Н	36.006	37.267	26.500	20.00
2001	TYR	HA	35.959	34.369	27.208	20.00
2002	TYR	1HB	38.106	35.477	26.945	20.00
2003	TYR	2HB	37.949	34.775	28.517	20.00
2004 2005	TYR	HD1	37.433	36.298 37.831	30.508 26.601	20.00 20.00
2005	TYR TYR	HD2 HE1	38.288 38.028	38.353	31.577	20.00
2007	TYR	HE2	38.875	39.932	27.553	20.00
2007	TYR	HH	37.890	41.054	30.183	20.00
2009	TRP	N	34.243	36.345	28.877	11.94
2010	TRP	CA	33.260	36.444	29.897	12.84
2011	TRP	C	31.795	36.329	29.296	14.59
2012	TRP	Ö	31.599	36.587	28.134	14.57
2013	TRP	CB .	33.665	37.712	30.717	12.85
2014	TRP	CG	33.087	38.918	30.092	13.65
2015	TRP	CD1	31.828	39.488	30.403	14.58
2016	TRP	CD2	33.657	39.662	29.006	12.43
2017	TRP	NE1	31.613	40.540	29.550	14.02
2018	TRP	CE2	32.726	40.705	28.701	12.63

2051 GLN       CB       27.240       38.783       26.497         2052 GLN       CG       28.579       39.367       26.064         2053 GLN       CD       28.474       39.304       24.551         2054 GLN       OE1       28.270       38.247       23.966         2055 GLN       NE2       28.566       40.493       23.988         2056 GLN       H       28.103       36.536       27.563         2057 GLN       HA       27.828       39.333       28.450         2058 GLN       1HB       26.462       39.506       26.222         2059 GLN       2HB       27.001       37.849       25.971         2060 GLN       1HG       29.420       38.737       26.370         2061 GLN       2HG       28.769       40.372       26.438         2062 GLN       1HE2       28.441       40.574       23.007         2063 GLN       2HE2       28.815       41.241       24.593         2064 LYS       N       25.408       37.425       28.909         2065 LYS       CA       24.255       37.495       29.783	32.71 36.57 35.28 40.87 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.39
2061 GLN       2HG       28.769       40.372       26.438         2062 GLN       1HE2       28.441       40.574       23.007         2063 GLN       2HE2       28.815       41.241       24.593	20.00 20.00 20.00 20.39 19.10

						40	4			
2071	LYS	CE	21.248		37.656		27.135		32.07	
2072	LYS	NZ	19.951		37.638		26.450		40.14	
2073	LYS	Н	25.913	,	36.568		28.818		20.00	
2074	LYS	HA	24.207		38.466		30.286		20.00	
2075	LYS	1HB	23.084		38.211		28.385		20.00	
2076	LYS	2HB	22.156		37.389		29.639		20.00	
2077	LYS	1HG	22.886		35.151		28.800		20.00	
2078	LYS	2HG	23.896		35.943		27.603		20.00	
2079	LYS	1HD	21.052		35.524		27.493		20.00	
2080	LYS	2HD	22.158		35.907		26.237		20.00	
2081	LYS	1HE	21.938		38.332		26.616		20.00	
2082	LYS	2HE	21.014		38.102		28.111		20.00	
2083	LYS	1HZ	19.364		37.019		27.058		20.00	
2084	LYS	2HZ	20.056		37.233		25.501		20.00	
2085	LYS	3HZ	19.574		38.608		26.433		20.00	
2086	GLU	N	23.400		36.776		31.841		15.88	
2087	GLU'	CA	23.136		36.118		33.111		15.37	
2088	GLU	C	22.807		34.620		32.974		15.69	
2089	GLU	0	23.432		33.804		33.626		13.68	
2090	GLU	CB	22.083		36.964		33.900		16.72	
2091	GLU	CG	22.625		38.280		34.481		14.29	
2092 2093	GLU GLU	CD OE1	22.320		39.420		33.579		18.80	
2093	GLU	OE2	22.205 22.216	-	39.180 40.551		32.359 34.077		21.62 21.27	
2095	GLU	H	22.210		37.629	,	31.683		20.00	
2096	GLU	HA	24.072		36.180		33.655		20.00	
2097	GLU	1HB	21.726		36.367		34.740		20.00	
2098	GLU	2HB	21.182		37.117		33.295		20.00	
2099	GLU	1HG	23.692		38.268		34.620		20.00	
2100	GLU	2HG	22.172		38.502		35.438		20.00	
2101	GLU	Ν	21.816		34.259	•	32.125		18.42	
2102	GLU	CA	21.276		32.842		31.945		20.95	
2103	GLU	С	22.156		31.928		31.090		20.76	
2104	GLÜ	0	21.994		30.705		31.057		20.49	
2105	GLU	СВ	19.952		32.834		31.234		20.78	
2106	GLU	CG	19.309		34.201		31.354		25.93	
2107	GLU	CD	19.642		35.143		30.246		25.99	
2108	GLU	OE1	19.653		34.697		29.122	•	28.63	
2109	GLU	OE2	19.906	4	36.310		30.524		25.53	
2110	GLU	H	21.335		35.045		31.723		20.00	
2111	GLU	HA	21.174		32.451		32.958		20.00	
2112	GLU	1HB	19.298		32.082		31.685	•	20.00	
2113	GLU	2HB	20.022		32.547		30.179		20.00	
2114 2115	GLU GLU	1HG 2HG	19.417 18.245		34.671		32.332		20.00	
2116	LYS	N	23.111		34.072 32.603		31.265 30.414		20.00	
2117	LYS	CA	24.125		31.842		29.692		21.38 21.50	
2118	LYS	C	25.526		31.935		30.345		21.97	
2119	LYS	0	26.347		32.738		29.892		22.57	-
2120	LYS	CB	24.088		32.258		28.172		22.92	
2121	LYS	CG	22.714		31.845		27.594		27.23	
2122	LYS	CD	22.552		31.376		26.144		35.59	
	-									

2123 2124 2125 2126 2127 2128 2129 2130 2131 2133 2133 2133 2133 2134 2143 2144 2145 2146 2153 2155 2156 2166 2167 2168 2168 2168 2168 2168 2168 2168 2168	LYSSSSSSSSSSSSUUUUUUUUUUUUUUUUUTTTTTTTTTT	H HA 1HB 2HB 1HG 2HG 1HE	23.728 24.007 23.100 23.905 24.907 24.269 21.975 22.353 22.496 21.598 24.630 23.573 23.210 24.158 24.854 25.773 27.139 28.092 27.261 26.205 25.370 25.536 24.429 27.261 28.206 25.454 26.595 27.261 28.206 25.454 26.595 29.391 30.416 31.427 30.870 32.137 33.223 29.589 29.978 32.321 31.744 33.506	30.467 29.243 33.602 30.779 31.743 33.331 32.616 30.996 32.263 30.867 31.088 30.193 28.748 31.078 30.999 30.064 29.223 30.453 29.525 28.477 27.271 28.906 31.330 30.212 29.090 30.275 29.603 28.585 29.473 30.660 31.714 32.886 31.855 30.862 29.130 30.175 31.163 31.258 31.214 31.511 30.974	25.664 26.485 30.448 29.750 27.668 28.039 27.813 28.180 25.506 26.26 26.26 26.26 26.26 27.490 27.490 26.137 31.083 33.369 33.369 33.224 33.369 33.224 33.488 32.545 31.701 31.877 34.014 33.571 34.324 34.874 31.312 30.511 31.349 32.433 30.153 29.625 29.762 31.066 29.604 28.283 26.943 28.435	43.39 46.48 20.00
2165 2166	MET MET	1HG 2HG	30.026 30.521	32.258 31.214	29.604 28.283	20.00
2168	MET	2HE	33.506	30.974	28.415	20.00
2169	MET	3HE N.	34.117 31.339	 32.425 27.348	27.610 30.934	 20.00 18.63
2171 2172	ILE	CA C	32.211 33.493	26.451 26.318	31.678 30.901	20.60 20.37
<ul><li>2173</li><li>2174</li></ul>	ILE	O CB	33.409 31.480	26.217 25.129	29.700 31.975	23.05 24.01

2175	ILE	CG1	30.423		25.362		33.119		25.81
2176	ILE	CG2	32.435	•	24.026		32.386		24.90
2177	ILE	CD1	29.015		25.092		32.627		27.88
2178	ILE	Н	30.974		27.074		30.050		20.00
2179	ILE	HA	32.485		26.885		32.636		20.00
2180	ILE	HB	30.985		24.804		31.061		20.00
2181	ILE	1HG1	30.463		26.402		33.452		20.00
2182	ILE	2HG1			24.754	•	34.001		20.00
2183	ILE		33.016		24.304		33.266		20.00
2184	ILE		33.118		23.779		31.571		20.00
2185	ILE		31.883		23.113		32.621	,	20.00
2186	ILE	1HD1	28.917		24.029		32.403		20.00
2187	ILE	2HD1			25.634		31.710		20.00
2188	ILE	3HD1	28.264		25.343		33.374		20.00
2189	PHE	N	34.683		26.352		31.568		19.97
2190	PHE	CA	35.937		26.121		30.909		18.46
2191	PHE	C	36.319		24.745	,	31.318		19.73
2192	PHE	Ō	36.707		24.506		32.444		18.94
2193	PHE	СВ	36.932		27.287		31.208		16.35
2194	PHE	CG	36.422		28.683		31.025		14.87
2195	PHE	CD1	36.522		29.299		29.779		13.55
2196	PHE	CD2	35.733		29.310		32.097		16.14
2197	PHE	CE1	35.830		30.461		29.569		14.72
2198	PHE	CE2	35.066		30.492		31.858		14.63
2199	PHE	CZ	35.099		31.052		30.579		14.92
2200	PHE	Н	34.651		26.525		32.549		20.00
2201	PHE	HA	35.764		26.097		29.830		20.00
2202	PHE	1HB	37.703		27.282		30.445		20.00
2203	PHE	2HB	37.090	•	27.353		32.285	,	20.00
2204	PHE	HD1	37.082		28.849		28.984		20.00
2205	PHE	HD2	35.695		28.842		33.075		20.00
2206	PHE	HE1	35.848		30.933	• •	28.597		20.00
2207	PHE	HE2	34.533		30.984		32.644		20.00
2208	PHE	ΗZ	34.566	٠,	31.975		30.381		20.00
2209	GLU	N	36.121		23.790		30.412		23.42
2210	GLU	CA	36.337		22.339		30.733		27.04
2211	GLU	C	37.729		21.979		30.888		26.67
2212	GLU	0	38.100		21.211		31.739		27.77
2213	GLU	CB	35.728		21.345		29.766		32.84
2214	GLU	CG	34.183		21.469		29.799		43.59
2215	GLU	CD	33.519		20.760		28.583		52.36
2216	GLU	OE1	33.847		21.115		27.436		54.69 55.07
2217	GLU	OE2	32.681		19.880		28.790 29.529	•	20.00
2218	GLU	Н	35.751		24.069 22.173		31.709		20.00
2219 2220	GLU GLU	HA 1HB	35.881 35.996		20.309		30.011		20.00
2221	GLU	2HB	36.098		21.541		28.755		20.00
2222	GLU	1HG	33.904		22.517		29.697		20.00
2223	GLU	2HG	33.744		21.100		30.727	- 11 Oc	20.00
2224	ASP	N N	38.560		22.594		30.081		25.27
2225	ASP	CA	39.984		22.349		30.299		24.77
2226	ASP	C	40.545		22.756		31.677		24.82

2227 2228 2229 2230 2231 2232 2233 2234 2235 2236 2237 2240 2241 2242 2243 2244 2245 2246 2247 2248 2249 2250 2251 2252 2253 2256 2257	ASP THR		41.386 40.679 40.734 39.805 41.734 38.274 40.119 40.199 41.699 40.124 40.690 39.585 39.768 41.135 40.007 42.544 39.685 41.531 41.412 39.736 42.651 43.224 42.971 38.400 37.484 36.941 36.879 38.125 37.063 36.007 37.398		22.063 23.052 24.550 25.153 25.112 23.318 21.276 22.776 22.679 23.834 24.144 23.979 24.233 25.633 26.430 25.763 24.464 23.510 25.917 26.169 25.408 25.917 26.169 25.408 25.917 26.169 25.408 25.917 26.169 25.408 25.751 25.408 26.751 27.769 27		32 189 29 135 29 259 29 754 28 924 29 453 30 225 28 195 29 067 32 305 33 672 34 752 35 885 33 587 33 110 32 851 31 679 33 937 34 651 32 225 31 820 33 381 32 905 34 414 35 497 36 323 37 538 36 450 37 882		26.69 25.87 28.11 27.95 30.09 20.00 20.00 20.00 20.10 20.15 18.17 15.16 16.50 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.10 20.00
	•				1				
		CB			22.180		36.450		27.97
			• •	.,				-	
						 		•	
225 <i>7</i> 2258	ASN ASN	ND2 H	37.398 38.197		20.416		37.882		20.00
2259	ASN	HA	36.637		22.771		34.958	•	20.00
2260	ASN	1HB	38.630		22.649		37.291		20.00
2261	ASN	2HB	38.876		21.572	,	35.956		20.00
2262	ASN	,	36.827		19.614		38.005		20.00
2263 2264	ASN LEU	2HD2 N	38.154 36.523		20.743 25.343		38.435 35.557		20.00 22.74
2265	LEU	CA	36.054		26.662		36.015		21.67
2266	LEU	C	34.666		26.976		35.433		20.94
2267	LEU	.0	34.360		26.609		34.348		21.39
2268	LEU	CB	37.062		27.693		35.512		21.84
2269 2270	LEU LEU	CG CD1	38.192 39.526		28.179 28.185		36.501 35.755		23.27 17.13
2270	LEU	CD1	38.304		27.532	•	37.894		21.62
2272	LEU	Н	36.596		25.109	*,	34.588		20.00
2273		HA	35.977		26.636	• •	37.101		20.00
2274	LEU	1HB	36.570		28.580		35.111		20.00
2275 2276	LEU LEU	2HB HG	37.540 37.973		27.261 29.219		34.632 36.721		20.00
2270	LEU	1HD1			28.735		34.816		20.00
2278	LEU	2HD1			27.168		35.542		20.00

2279	LEU	3HD1	40.281		28.627		36.394		20.00
2280	LEU		38.454		26.455		37.798		20.00
2281	LEU		37.409				38.497		20.00
					27.696				
2282	LEU	3HD2			27.916		38.459		20.00
2283	LYS	N	33.788		27.630		36.164		21.10
2284	LYS	CA	32.465		28.108		35.683		19.53
2285	LYS	С	32.490		29.604		35.894		18.34
2286	LYS	0	33.087		30.091		36.832		20.22
2287	LYS	CB	31.333		27.455		36.480		20.94
2288	LYS	CG	29.922		27.807		36.039		22.87
2289	LYS	CD.	28.975		26.839		36.717		29.27
2290	LYS	CE -	27.555		27.322		37.117		32.16
2291	LYS	NZ	27.241		26.588		38.379		36.84
2292	LYS	H	34.067		27.812		37.110		20.00
2293	LYS	HA	32.356		27.890		34.626		20.00
2294	LYS	1HB	31.418	٠	27.687	,	37.535		20.00
2295	LYS	2HB	31.452		26.379		36.378		20.00
2296	LYS	1HG	29.776		27.833		34.960		20.00
2297	LYS	2HG	29.738		28.789		36.395		20.00
2298	LYS	1HD	29.460		26.460		37.618		20.00
2299	LYS	2HD	28.872		25.967		36.075		20.00
2300	LYS	1HE	26.811		27.141		36.334		20.00
2301	LYS	2HE	27.560		28.394		37.332		20.00
2302	LYS	1HZ	28.022		26.733		39.071		20.00
2302	LYS	2HZ	27.136		25.563		38.252		20.00
	LYS	3HZ			26.914		38.881		20.00
2304			26.383				35.029		16.41
2305	LEU	N	31.850		30.299				
2306	LEU	CA	31.816		31.744		35.238		16.41
2307	LEU	С	30.414	*	32.310		34.879		16.89
2308	LEU	Ο.	29.861.		31.985		33.848		16.71
2309	LEU	CB	32.856		32.320		34.286		15.09
2310	LEU	CG	33.093		33.868		34.349	•	13.98
2311	LEU	CD1	33.803		34.225		33.001		15.42
2312	LEU	CD2	33.869	•	34.360		35.697		16.75
2313	LEU	Н	31.410		29.830		34.259		20.00
2314	LEU	HA	32.044		32.013		36.257		20.00
2315	LEU	1HB	32.580		32.030		33.270		20.00
2316	LEU	2HB	33.813		31.832		34.474		20.00
2317	LEU	HG	32.130	•	34.370		34.324		20.00
	LEU	1HD1			33.816		32.134		20.00
	LEU		34.821		33.849		32.995		20.00
	LEU		33.860		35.305		32.873		20.00
2321	LEU		34.834		33.868		35.776		20.00
2322	LEU		33.302		34.161		36.605		20.00
2323	LEU		34.065		35.433		35.659		20.00
2324	THR	N	29.882		33.181	- 	35.720		17.65
2325	THR	CA	28.551		33.725		35.460		17.51
	THR-		28.621		35.167		35.480		16.95
2327	THR	0	29.498		35.674		36.178		16.10
2328	THR	CB	27.606		33.300		36.590		15.94
2329	THR	OG1	27.778		31.872		36.920		15.73
2330	THR	CG2	26.172		33.301		36.033		17.67

2383 2384 2385 2386 2387 2388 2390 2391 2392 2393 2394 2395 2396 2397 2398 2399 2400 2401 2402 2403 2404 2405 2406 2407 2408 2409	SERRUUUUUUUUUUUUUUUUUPPPPPPPPPPPPPPPPPPP	HABB HACCOCCCOOH HABBGG HABGG HABBGG HABGG HABBGG HABG HAB	24.533 24.803 24.053 26.904 25.933 26.837 26.217 25.011 27.228 26.075 26.377 27.547 25.405 24.982 27.770 27.987 27.650 25.222 25.745 27.039 26.705 27.443 28.624 27.189 27.254 26.258 28.321	41.708 42.025 43.172 43.995 43.809 44.927 46.306 46.358 44.878 44.442 44.181 44.144 43.982 43.809 44.930 44.106 45.808 45.118 43.467 47.342 48.761 49.821 49.821 49.177 50.675 51.344 51.154	36.356 39.038 38.066 38.253 35.222 34.671 34.923 34.981 33.158 32.355 30.922 30.531 30.205 34.916 35.236 33.047 32.773 32.423 32.773 32.423 32.706 35.079 35.113 33.921 33.900 36.507 36.768 36.462 37.274	20.00 20.00 20.00 20.00 22.91 25.57 25.11 28.12 27.94 32.29 38.30 43.02 41.87 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.31 18.75 19.30 17.85 21.92 28.13 28.01 33.77
<ul><li>2414</li><li>2415</li><li>2416</li></ul>	ILE	N	26.646	49.500	32.878	19.47
	ILE	CA	27.011	50.169	31.640	20.98
	ILE	C	26.884	51.689	31.756	22.51
<ul><li>2417</li><li>2418</li><li>2419</li></ul>	ILE	O	25.854	52.187	32.173	25.15
	ILE	CB	26.054	49.679	30.540	22.51
	ILE	CG1	26.166	48.190	30.336	22.32
2420	ILE	CG2	26.210°	50.376	29.202	23.73
2421	ILE	CD1	25.344	47.693	29.157	22.61
2422 · 2423 · 2424	ILE ILE	HA HB	25.726 28.044 25.034	49.152 49.924 49.883	33.013 31.444 30.893	20.00 20.00 20.00
2425 2426 2427	ILE	2HG1	25.831 27.204 26.990	47.659 47.891 49.941	31.231 30.185 28.588	20.00 20.00 20.00
2428	ILE	2HG2	26.412	51.439	29.278	20.00
2429	ILE	3HG2	25.287	50.309	28.630	20.00
<ul><li>2430</li><li>2431</li><li>2432</li></ul>	ILE ILE ILE	2HD1	25.810 24.316 25.309	47.995 48.054 46.600	28.217 29.173 29.147	20.00 20.00 20.00
	LYS	N	27.927	52.396	31.334	21.11
	LYS	CA	28.007	53.844	31.411	20.45

2435 2436 2437 2438 2443 2443 2444 2444 2445 2445 2453 2454 2453 2454 2453 2454 2463 2464 2467 2473 2474 2475 2476 2477 2478 2478 2479 2479 2479 2479 2479 2479 2479 2479	SSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSS	2HG2	28.454 28.625 29.094 28.926 28.269 27.931 27.390 28.708 27.026 29.330 30.013 29.915 28.387 27.317 28.840 28.821 27.183 26.728 28.153 26.891 28.665 28.839 30.207 30.252 28.731 29.994 27.406 28.394 28.017 28.524 30.033 27.363 26.606 27.015 31.305 32.678 33.364 33.967 33.364 35.206 33.908 35.785 35.129 35.664	54.260 53.399 54.210 53.803 54.939 54.725 56.048 51.893 54.274 55.273 53.731 53.631 52.862 55.140 55.855 54.455 53.930 56.433 56.770 56.014 55.518 55.342 57.347 57.972 57.828 56.233 57.312 57.312 57.312 57.312 57.313 57.757	29.997 29.162 32.440 33.916 34.702 36.183 36.625 30.963 31.645 32.393 32.121 34.341 34.018 34.208 34.583 36.758 36.298 35.914 36.673 37.524 29.676 28.253 27.586 26.399 28.048 28.523 27.586 26.399 28.048 28.523 28.730 30.314 27.719 26.923 28.697 29.806 28.421 28.446 27.957 28.591 28.162 28.157 29.464 30.382 29.786 31.591 30.989 31.908 33.129	20.80 22.26 22.37 25.62 29.83 33.05 35.32 20.00
2478	TYR	CE2	35.785	56.566	30.989	16.69
					•	
2481	TYR.	H :	31.090	55.949	29.230	20.00
2482 2483	TYR TYR	HA 1'HB	32.616 32.511	54.883 57.176	26.881 28.055	20.00
2484	TYR	2HB	33.813	56.779	27.310	20.00
2485 2486	TYR TYR	HD1 HD2	32.439 35.662	58.052 55.472	30.168 29.127	20.00 20.00

2487	TYR	HE1	33.342	58.642	32.206	20.00
2488	TYR	HE2	36.782	56.174	31.160	20.00
2489	TYR	HH	35.472	57.124		
					33.840	20.00
2490	TYR	N	32.720	53.248	29.650	14.74
2491	TYR	CA	33.167	52.050	30.352	14.03
2492	TYR	С	31.906	51.359	30.862	16.12
2493	TYR	0	30.824	51.930	30.915	17.82
2494	TYR	СВ	34.251			
				52.412	31.375	12.74
2495	TYR	CG	33.728	53.134	32.579	15.40
2496	TYR	CD1	33.475	52.431	33.758	13.77
2497	TYR	CD2	33.476	54.490	32.544	18.58
2498	TYR	CE1	33.015	53.078	34.883	15.81
2499	TYR	CE2	32.944	55.161	33.649	17.84
2500	TYR	CZ	32.740		*	
				54.435	34.866	16.69
2501	TYR	ОН	32.240	54.871	36.111	. 18.92
2502	TYR	Н	31.850	53.689	29.886	20.00
2503	TYR	HA	33.628	51.368	29.650	20.00
2504	TYR	1HB	35.016	53.026	30.901	20.00
2505	TYR	2HB	34.748	51.510	31.713	20.00
2506	TYR	HD1	33.656	51.367		
					33.806	20.00
2507	TYR	HD2	33.632	55.018	31.619	20.00
2508	TYR	HE1	32.854	52.539	35.807	20.00
2509	TYR	HE2	32.615	56.155	33.358	20.00
2510	TYR	HH	31.864	55.766	36.179	20.00
2511	THR	N	32.096	50.092	31.166	14.21
2512	THR	CA	31.125	49.193	31.834	15.42
2513						
	THR	C	31.889	48.540	33.043	15.44
2514	THR	0	33.128	48.344	32.955	` 15.14
2515	THR	CB	30.572	48.077	30.896	16.15
2516	THR	OG1	29.894	48.442	29.632	16.75
2517	THR	CG2	29.525	47.168	31.562	15.21
2518	THR	Н	33.038	49.794	31.028	20.00
2519	THR	HA	30.320	49.806	32.254	20.00
2520						
	THR	HB	31.408	47.328	30.806	20.00
2521	THR	HG1	30.097	49.274	29.118	20.00
2522	THR	1HG2	28.699	47.761	31.955	20.00
2523	THR	2HG2	29.934	46.590	32.391	20.00
2524	THR	3HG2	29.096	46.451	30.861	20.00
2525	VAL	N	31.079	48.256	34.137	14.25
2526	VAL	CA	31.584	47.499		
2527	VAL				35.318	15.61
		C	30.715	46.291	35.625	15.76
2528	VAL	0	29.577	46.421	36.050	16.87
	VAL ,	CB	31.794	48.216	36.666	15.56
2530	VAL <sup>2</sup>	CG1	32.554	49.567	36.570	17.93
2531	VAL	CG2	32.587	47.191	37.535	15.84
2532	VAL	Н	30.179	48.701	34.161	20.00
2533	VAL	HA ·	32.559	47.126	35.001	
						20.00
2534	VAL	HB	30.835	48.422	37.142	20.00
2535			31.976	50.307	36.013	20.00
2536	VAL		33.491	49.416	36.039	20.00
2537	VAL	3HG1	32.779	49.960	37.562	20.00
2538	VAL	1HG2	33.506	46.870	37.041	20.00
	_			<b>.</b>	- · · · · · ·	_0.00

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2695	THR	2HG2	2 39.155		20.320	•	43.413		20.00
2696	THR		2 40.498		20.158		42.403		20.00
2697	THR	N	36.755						43.51
					22.357		43.620		
2698	THR	CA	36.569		22.492		45.080		40.76
2699	THR	C ·	35.225		23.028		45.501		41.34
2700	THR	0	34.949	•	23.173		46.685		43.02
2701	THR	CB	37.422		23.692		45.561		39.25
2702	THR	OG1	37.152		25.022		44.987		42.09
2703	THR	CG2	38.908		23.417		45.377		37.92
2704	THR	Н	36.887		23.135		43.004		20.00
2705	THR	HA	36.765		21.561	•	45.612		20.00
2706	THR	HB	37.364		23.699		46.689		20.00
2707	THR	HG1	36.526		25.122		44.220		20.00
2708	THR	1HG2			23.284		44.335		20.00
2709	THR	2HG2			22.506		45.911		20.00
2710	THR		39.483		24.240		45.802		20.00
2711	GLN	N	34.476		23.431		44.476		40.94
2712	GLN	CA	33.352		24.345		44.557		41.20
2713	GLN	C	33.545		25.695		45.299		38.70
2714	GLN	Ö	32.586	- 111	26.414		45.546	•	38.13
2715	GLN	СВ	32.149						
2716	GLN				23.549		45.037		45.37
		CG -	31.786	, -	22.523		43.980		51.51
2717	GLN	CD	30.414		21.959		44.267		56.44
2718	GLN	OE1	30.270		21.226		45.234		58.04
2719	GLN	NE2	29.438		22.272		43.407	•	61.09
2720	GLN	Н	34.741		23.135	-	43.558		20.00
2721	GLN	HA	33.160		24.656		43,530		20.00
2722	GLN	1HB	31.310		24.234		45.188		20.00
2723	GLN	2HB	32.358		23.063		45.989		20.00
2724	GLN	1HG	32.495		21.702		43.926		20.00
2725	GLN	2HG	31.753		22.994		42.994		20.00
2726	GLN	1HE2	28.585		21.754		43.419		20.00
2727	GLN	2HE2	29.601	00	22.971°		:42.703		20.00
2728	GLU	N	34.762		26.092		45.656		37.29
2729	GLU	CA	34.911		27.514		46.023		37.74
2730	GLU	С	34.377		28.454		44.928		35.73
2731	GLU	0	34.427		28.209		43.728		34.94
2732	GLU	СВ	36.317		27.954	•	46.489		43.18
2733	GLU	CG	36.377		29.427		47.019		51.01
2734	GLU	CD	37.799		29.866		47.427		57.11
2735	GLU	OE1	38.637		29.023		47.594		59.70
2736	GLU	OE2	38.063	·	31.062		47.546		59.23
2737	GLU	H	35.487		25.418		45.773		20.00
2738	GLU	HA	34.261		27.618	•	46.889	•	20.00
2739		1HB	37.020	- 1	27.876		45.674		20.00
2740	GLU	2HB	36.685		27.267		47.255		20.00
2741	GLU	1HG	35.747 <sup>4</sup>		29.531·	я · ·	47.902		20.00
2742	GLU	2HG	36.060		30.154		46.279		20.00
2743	THR	N	33.861		29.566		45.444	<del></del>	33.44
2744	THR	CA	33.250		30.618		44.592		31.67
2745	THR	CA	33.937		31.934	2	44.874	٠	
2745									30.34
2140	THR	0	34.285		32.227		46.011		31.62

2747	THR	СВ	31.701	30	.674	44	.951		<sup></sup> 29.52
2748	THR	OG1	31.012		.973		.882		31.41
2749	THR	CG2	31.094		.901		.741		27.83
2750	THR	H	34.048						
2751					.665		.415		20.00
	THR	HA	33.482		.372		.553		20.00
2752	THR	HB	31.695		.004		.845		20.00
2753	THR	HG1	30.437		.433		.203		20.00
2754	THR		2.31.122		.846		.187		20.00
2755	THR		2 31.678		.101		635		20.00
2756	THR		2 30.068		.750	,	.091		20.00
2757	ARG	Ν	34.107	32.	738	43.	860		27.40
2758	ARG	CA	34.538	34.	.090	44.	139		22.96
2759	ARG	С	33.959	35.	.067	43.	175		19.60
2760	ARG	Ο .	33.722	34.	760	42.	031		18.08
2761	ARG	CB	36.078	34.	229	44.	003		24.02
2762	ARG	ÇG	36.854	33.	227	44.	797		27.06
2763	ARG	CD	38.285	33.	547	44.	508		30.34
2764	ARG	NE	39.183	32.	683	45.	219		31.08
2765	ARG	CZ	40.370	32.	287	44.	813	*	33.27
2766	ARG	NH1	41.054	32.	842	43.	773		28.01
2767	ARG	NH2	40.742	31.	202		477		34.77
2768	ARG	Н	33.782		444		964		20.00
2769	ARG	HA	34.184		369		135		20.00
2770	ARG	1HB	36.387		247		261		20.00
2771	ARG	2HB	36.342		082		952		20.00
2772		1HG	36.606		197		535		20.00
277.3	ARG	2HG	36.637		357		857		20.00
2774	ARG	1HD	38.521		562		846		20.00
2775	ARG	2HD	38.485		476		435		20.00
2776	ARG	HE	38.854		134		993		20.00
2777	ARG	1HH1	41.905		439		426		20.00
2778	ARG	2HH1	40.666		647		339		20.00
2779	ARG		41.602	30.		45.			20.00
2780	ARG		40.159		814	46.2			20.00
2781	GLU	N	33.859		255	43.			19.61
2782	GLU	CA	33.468	37.4		43.0			20.49
2783	GLU	C	34.699	38.2		42.			18.35
2784	GLU	Ö	35.609	38.6		43.			19.50
2785	GLU	СВ	32.651	38.		43.9			23.73
2786	GLU	CG	32.354	39.6		43.			30.58
2787	GLU	CD	31.697	40.6		44.(			37.37
2788	GLU	OE1	32.349	41.6		44.0			
2789	GLU	OE2	30.572	40.4					42.91
2790	GLU	H	34.048			44.4			39.95
2791	GLU	HA	32.845	36.2		44.7			20.00
2792				37.		42.2			20.00
2792 2793 .	GLU GLU	1HB 2HB	33.179	38.6		44.8			20.00
2793 . 2794	GLU		31.718	37.9		44.2			20.00
2795	GLU	1HG	31.671	39.4		42.2			20.00
2795		ZHG	33.211	40.1		42.6			20.00
	ILE	N	34.562	38.4		41.2			17.31
2797	ILE	CA	35.533	39.0		40.4			15.44
2798	ILE	C	34.806	40.3	529	39.8	120		13.03

2799 2800 2801 2802 2803 2804 2805 2806 2807 2808 2810 2811 2812 2813 2814 2815 2816 2817 2818 2819 2820 2821 2822 2823 2824	ILEEEEEEEEEEEUUUUUUUUUUUUUUUUUUUUUUUUUU	1HG2 2HG2	33.725 36.014 36.469 37.156 37.777 33.698 36.373 35.178 36.634 35.686 37.976 36.753 37.571 37.651 38.583 38.126 35.437 34.934 35.639 36.823 35.395 34.233 34.708 33.503 36.344 33.859		40.269 38.081 36.717 38.781 36.731 38.103 39.393 37.891 36.068 36.219 39.123 39.664 38.142 37.273 37.221 35.731 41.444 42.684 42.846 42.846 43.657 43.955 44.759 42.721 41.402 42.644		39.307 39.316 39.808 38.511 40.616 40.833 41.053 38.645 38.940 40.382 39.142 38.013 37.731 41.553 40.068 40.878 39.855 39.281 37.908 37.768 40.382 41.312 42.493 41.757 40.277 39.125		13.73 14.77 13.99 13.81 13.39 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 9.92 10.85 12.45 13.72 11.52 17.76 14.86 15.60 20.00 20.00
2809	ILE	1HG2	37.976		39.123		39.142		20.00
				,		•			
2811	IĿĘ	3HG2	37.571		38.142				
	-								
				`					
		_							
2825	LEU	1HB	35.739		44.594		39.965		20.00
2826	LEU	2HB	36.256		43.272		40.931		20.00
2827	LEU	HG	33.518		44.568		40.763		20.00
2828	LEU	1HD1	35.021		45.750		42.182		20.00
2829	LEU	2HD1	35.528		44.252		42.999		20.00
2830	LEU	3HD1	33.915		44.895		43.226		20.00
2831	LEU		34.144		41.977		42.215		20.00
2832	LEU		32.928		42.261		40.949		20.00
2833	LEU	3HD2	32.747		43.012		42.487		20.00
2834	HIS	N	34.933		43.394		36.894		10.68
2835	HIS	CA	35.545		43.475		35.600		11.55
2836	HIS	C	35.323		44.872		35.144		12.79
2837	HIS	0	34.194		45.266		34.919	,	14.70
2838	HIS	СВ	34.788		42.489		34.668		11.04
2839	HIS	CG	35.263		42.384		33.227		9.68
2840	HIS	ND1	34.526		42.828		32.187		10.25
2841	HIS	CD2	36.429		41.786		32.744		10.64
2842	HIS	CE1 NE2	35.183 36.329		42.511 41.906		31.063 31.409		12.62 11.21
2843 2844	HIS HIS	H	33.982		43.625		37.069		20.00
2845		HA	36.601		43.242		35.581		20.00
2846	HIS	1HB	33.743		42.742		34.647		20.00
-2847 <sup></sup>		2HB	34.856		41.492		35.083		20.00
2848	HIS	HD1	33.677		43.272		32.321		20.00
2849	HIS	HD2	37.206		41.325		33.337		20.00
2850	HIS	HE1	34.876		42.676		30.032		20.00

2903 2904 2905 2906 2907 2908 2910 2911 2912 2913 2914 2915 2916 2917 2918 2919 2921 2921 2921 2921 2921 2921	TRP	2HG2 3HG2 N CA C O CB 1 CB 1HG2 2HG2 3HG2 N CA C O CB CD1 CE3 CC CC	33.561 33.378 38.641 40.120 40.640 40.679 40.854 42.300 43.194 43.010 44.399 44.332 42.647 45.251 43.596 44.901 38.223	53.598 52.873 51.639 53.065 53.478 53.926 53.434 54.166 53.890 53.113 54.034 53.268 53.730 52.591	30.426 27.111 31.176 26.713 30.901 28.064 27.075 25.758 24.708 23.530 25.763 25.886 26.638 27.614 25.389 24.715 26.739 27.675 26.159 25.104 24.097 23.978 23.501 24.407 25.757 24.541 26.052 23.110 23.415 25.953 25.327 23.626 24.814 24.218 22.987 22.980 25.467 25.605 24.620 26.877 25.165 26.576 28.170 27.615 29.207 28.923 24.823	20.00 20.00 20.00 20.00 20.00 13.66 14.87 16.37 19.74 13.92 14.21 14.13 20.00
						20.00
2952 2953	TRP TRP	HA 1HB	40.290 40.803	54.667 51.977	24.218 25.516	20.00 20.00
2954	TRP	2HB	40.369	53.437	26.358	20.00

2955	TRP	HD1	42.974		54.062		23.567		20.00
2956		HE1	45.198		54.476		24.680		20.00
2957		HE3	41.644		52.763		28.381		
									20.00
2958		HZ2	46.233		54.366		27.330		20.00
2959	TRP	HZ3	43.327		53.028		30.234		20.00
2960	TRP		45.624		53.867		29.709		20.00
2961	PRO	N	41.054		53.592		21.917	'	11.63
2962	PRO	CA	41.341		52.864		20.657		12.03
2963	PRO	С	42.762		52.223		20.620		10.91
2964	PRO	0	43.674		52.589		21.352		11.74
2965	PRO	СВ	41.002		53.928		19.599		15.45
2966	PRO	CG	40.318		55.060		20.382		16.92
2967	PRO	CD	40.930		55.068				
2968	PRO	HA					21.726		12.40
			40.641		52.036		20.543		20.00
2969	PRO	1HB	40.376	٠	53.540		18.790		20.00
2970	PRO	2HB	41.911		54.325		19.146		20.00
2971	PRO	1HG	40.429		56.033	•	19.909		20.00
2972	PRO	2HG	39.258		54.863		20.489		20.00
2973	PRO	1HD	40.296		55,584		22.434		20.00
2974	PRO	2HD	41.916		55.543		21.732		20.00
2975	ASP	Ν	42.866		51.223		19.711		9.83
2976	ASP	CA	44.171		50.530		19.648		13.50
2977	ASP	С	45.160	1	51.471		19.077		15.42
2978	ASP	Ö	44.802		52.201		18.172	٠.	16.15
2979	ASP	СВ	44.065		49.223		18.819		14.19
2980	ASP	CG.	45.153		48.241		19.221		17.10
2981	ASP	OD1	45.877		48.540		20.150		16.38
2982	ASP	OD2	45.270	, .	47.155		18.676		16.84
2983	ASP	H	42.077		50.813		19.277		20.00
2984	ASP	HA	44.449		50.328		20.691		20.00
2985	ASP	1HB							
			44.135		49.419		17.751		20.00
2986	ASP	2HB	43.120	*	48.725		19.033		20.00
2987	PHE	Ň	46.329		51.527		19.651		15.66
2988	PHE	CA	47.349		52.538		19.450		13.22
2989	PHE	C	46.921		54.020		19.555		13.65
2990	PHE	. 0	47.504		54.923		18.956		14.13
2991	PHE	CB	48.233		52.272		18.290		15.03
2992		CG	48.749		50.895		18.233		16.37
2993	PHE	CD1	49.950		50.591		18.833		12.85
2994	PHE	CD2	48.009		49.927		17.529		15.04
2995	PHE	CE1	50.376	•	49.294		18.803	,	10:54
2996	PHE	CE2	48.460		48.659		17.407		12.82
2997	PHE	CZ	49.626		48.341		18.105		12.68
2998	PHE	Н	46.546		50.666		20.113		20.00
2999	PHE	HA	47.975		52.405	•	20.321	:	20.00
3000	PHE	1HB	49.022		53.011		18.302		20.00
3001	PHE	2HB	47.662		52.452		17.380		20.00
3002	PHE	HD1	50.490		51.363		19.335		20.00
3002		пD1 -HD2-	47.070		50.181		19.335		
3003		HE1							20.00
	PHE		51.281		48.969		19.299		20.00
	PHE	HE2	47.901		47.906		16.873		20.00
3006	PHE	HZ	49:963		47.312		18.117		20.00

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3007 3008 3009 3010 3011 3012	GLY GLY GLY GLY	N CA C O H 1HA	45.887 45.240 45.436 46.132 45.489 44.176		54.153 55.436 55.859 55.250 53.315 55.267	20.424 20.680 22.144 22.953 20.797 20.536		12.17 12.74 13.53 12.80 20.00 20.00	
3013 3014 3015 3016 3017 3018	GLY VAL VAL VAL VAL	2HA N CA C O CB	45.596 44.801 44.812 43.249 42.426 45.673		56.211 56.938 57.503 57.513 57.380 58.788	19.988 22.435 23.722 24.071 23.136 23.744		20.00 14.29 13.36 15.85 16.84 12.64	
3019 3020 3021 3022 3023 3024	VAL VAL VAL VAL VAL	CG1 CG2 H HA HB 1HG1	45.201 47.178 44.211 45.291 45.561 44.171		59.840 58.444 57.343 56.801 59.232 60.133	22.763 23.585 21.732 24.402 24.737 22.945		13.96 12.56 20.00 20.00 20.00 20.00	
3025 3026 3027 3028 3029 3030	VAL VAL VAL VAL PRO	3HG1 1HG2 2HG2	45.278 45.807 47.395 47.478 47.782 42.963		59.537 60.743 58.056 57.697 59.337 57.517	21.722 22.867 22.589 24.315 23.714 25.476		20.00 20.00 20.00 20.00 20.00 16.09	
3031 3032 3033 3034 3035 3036	PRO PRO PRO PRO PRO PRO	CA C O CB CG	41.668 41.138 41.887 42.035 43.260 44.019		57.936 59.345 60.270 58.106 57.208 57.339	26.088 25.657 25.348 27.586 27.767 26.463		14.29 16.70 15.47 12.24 11.83 14.99	
3037 3038 3039 3040 3041 3042	PRO PRO PRO PRO PRO	HA 1HB 2HB 1HG 2HG	40.885 41.236 42.288 43.844 42.915	.•	57.194 57.797 59.134 57.437 56.182	25.936 28.246 27.841 28.646 27.868		20.00 20.00 20.00 20.00 20.00	
3043 3044 3045 3046 3047	PRO PRO GLU GLU GLU	CA C O	44.640 44.642 39.830 39.149 39.701 39.946	٠.	56.472 58.226 59.499 60.735 61.877 62.949	26.300 26.489 25.747 25.401 26.121 25.567		20.00 20.00 18.37 20.99 22.01 24.26	
3048 3049 3050 3051 3052 3053	GLU GLU GLU GLU GLU	CB CG CD OE1 OE2 H	37.652 37.095 36.818 37.792 35.603 39.307	. 4	60.699 59.481 58.252 57.706 57.899 58.672	25.572 24.814 25.737 26.363 25.789 25.934		22.54 31.50 36.66 31.28 36.17 20.00	
3054 3055 3056 3057 3058	GLU GLU GLU GLU GLU	HA 1HB 2HB 1HG 2HG	39.384 37.225 37.322 37.751 36.147		60.894 61.596 60.660 59.146 59.734	 24.354 25.130 26.604 24.012 24.345	<del>-</del> -	20.00 20.00 20.00 20.00 20.00	

3059	SER	Ν	39.960	61.603	27.399	20.38
3060	SER	CA	40.647	62.714	28.080	17.88
3061	SER	С	41.167	62.167	29.306	15.08
3062	SER	0	40.789	61.054	29.671	14.67
3063	SER	CB	39.666	63.853	28.431	16.14
3064	SER	OG	38.604	63.400	29.339	13.25
3065	SER	Н	39.673	60.707	27.729	20.00
3066	SER	HA	41.461	63.055	27.454	20.00
3067	SER	1HB	39.386	64.462	27.511	20.00
3068	SER	2HB	40.217	64.669	28.926	20.00
3069	SER	HG	38.043	62.566	29.237	20.00
3070	PRO	N	42.046	62.978	29.961	17.02
3071	PRO	CA	42.562	62.555	31.277	16.44
3072	PRO	С	41.527	62.381	32.332	16.53
3073	PRO	0	41.596	61.476	33.162	15.74
3074	PRO	CB	43.653	63.578	31.636 30.294	16.53 18.72
3075 3076	PRO PRO	.CG CD	43.993 42.716	64.277 64.193	30.294 29.431	16.72
3076	PRO	HA	43.044	61.584	31.179	20.00
3078	PRO	пА. 1HB	44.530	63.132	32.109	20.00
3079	PRO	2HB	43.256	64.327	32.324	20.00
3080	PRO	1HG	44.374	65.291	30.432	20.00
3081	PRO	2HG	44.794	63.718	29.803	20.00
3082	PRO	1HD	42.963	64.147	28.370	20.00
3083	PRO	2HD	42.115	65.075	29.631	20.00
3084	ALA	N	40.466	63.193	32.219	16.66
3085	ALA	CA	39.366	63.113	33.208	16.12
3086	ALA	C	38.576	61.824	33.103	15.23
3087	ALA	Ö	38.319	61.127	34.062	15.53
3088	ALA	СВ	38.449	64.319	32.990	15.29
3089	ALA	Н	40.478	63.899	31.513	20.00
3090	ALA	HA	39.809	63.153	34.207	20.00
3091	ALA	1HB	38.110	64.430	31.968	20.00
3092	ALA	2HB	38.976	65.236	33.241	20.00
3093	ALA	3HB	37.586	64.273	33.637	20.00
3094	SER	Ν	38.207	61.442	31.920	15.32
3095	SER	CA	37.563	60.086	31.870	15.79
3096	SER	С	38.482	58.892	32.167	13.45
3097	SER	Ο.	38.104	57.921	32.825	13.08
3098	SER	CB	37.106	59.914	30.447	17.95
3099	SER	OG	38.030	60.657	29.618	26.88
3100	SER	Н	38.374	62.009	31.111	20.00
3101	SER	HA	36.720	60.060	32.554	20.00
3102	SER	1HB	36.110	60.396	30.400	20.00
3103	SER	2HB	36.640	58.920	30.206	20.00
3104	SER	HG	38.996	60.405	29.428	20.00
3105	PHE	N	39.774	59.018	31.727	13.85
3106	PHE	CA.	40.788	57.998	32.090	12.76
3107	PHE	С	40.881	57.892	33.605	12.10
3108	PHE	O	40.898	56.801 58.305	34.114	14.79
3109	PHE	CB	42.188	58.305 57.467	31.403 32.071	14.01 12.89
3110	PHE	CG	43.256	57.467	32.07	12.03

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3215	PHE	HE2	41.841	59.192	43.957	20.00
3216	PHE	HZ	43.888	59.517	42.548	20.00
3217	LYS	, <b>N</b>	37.472	54.865	39.696	14.03
3218	LYS	CA	36.583	53.707	39.921	13.68
3219	LYS	C	37.382	52.381	40.226	13.76
3220	LYS	0	37.060	51.643	41.128	13.17
3221	LYS	CB	35.710	53.494	38.629	17.74
3222 3223	LYS LYS	CG CD	34.241 33.252	53.905 52.778	38.681 39.167	26.69 32.82
3224	LYS	CE	31.708	53.171	39.423	34.45
3225	LYS	NZ	30.586	52.149	39.327	39.18
3226	LYS	H	37.491	55.414	38.855	20.00
3227	LYS	HA	35.958	53.951	40.781	20.00
3228	LYS	1HB	35.782	52.475	38.241	20.00
3229	LYS	2HB	36.164	54.118	37.857	20.00
3230	LYS	1HG	34.004	54.112	37.648	20.00
3231	LYS	2HG	34.101	54.839	39.226	20.00
3232	LYS	1HD	33.656	52.326	40.073	20.00
3233	LYS	2HD	33.287	51.993	38.414	20.00
3234	LYS	1HE	31.446	53.975	38.749	20.00
3235	LYS	2HE	31.667	53.627	40.417	20.00
3236	LYS	1HZ	30.720	51.336	39.961	20.00
3237	LYS	2HZ	30.445	51.758	38.364	20.00
3238	LYS	3HZ	29.655	52.552	39.548	20.00
3239 3240	VAL VAL	N CA	38.441 39.117	52.045 50.767	39.435 39.646	13.89 11.86
3241	VAL	CA	39.668	50.735	41.109	11.86
3242	VAL	Ö	39.382	49.808	41.856	11.98
3243	VAL	CB	40.222	50.455	38.595	12.00
3244	VAL	CG1	39.714	50.431	37.083	11.88
3245	VAL	CG2	40.882	49.143	38.956	12.37
3246	VAL	Н	38.644	52.666	38.678	20.00
3247	VAL	HA	38.347	50.001	39.562	20.00
	VAL .		40.980	51.236	38.654	20.00
•	VAL		39.334	51,404	` 36.781	20.00
3250	VAL		38.911	49.717	36.936	20.00
3251	VAL		40.521	50.193	36.388	20.00
3252	VAL		40.141 41.338	48.344	38.985 39.940	20.00 20.00
	VAL VAL		41.657	49.168 48.868	38.246	20.00
	ARG	N	40.409	51.870	41.447	12.93
3256	ARG	CA	40.873	52.261	42.832	14.48
3257.		C	39.825	52.066	43.972	16.45
3258		Ŏ	39.966°	51.141	44.744	17.22
	ARG	ČВ	41.467	53.625	42.820	13.76
3260	ARG		42.889	53.536	42.265	12.22
3261.	ARG	CD	43.652	54.811	42.463	13.67
3262	ARG	NE	45.033	54.629	42.060	15.68
3263		CZ-	46.003	55.526	42.281	14.21
3264	ARG	NH1	45.760	56.603	42.913	12.91
3265	ARG		47.224	55.411	41.884	12.03
3266	ARG	Н	40.563	52.500	40.699	20.00

3306 3307 3308 3309 3310 3311 3312 3313 3314 3315 3316 3317	SER SER SER GLY GLY GLY GLY SER SER SER SER SER	N C C O C B G H A B B B B B C C O C B G C C C C C C C C C C C C C C C C C	37.684 37.252 36.916 36.444 36.799 37.027 36.553 37.678 38.672 38.104 35.743 35.911 36.049 37.756 37.245 36.586 37.276 36.752 36.543 37.770 37.593 35.572 36.155 35.639 38.700 38.524 40.089 40.792 38.903 38.508 39.722 40.726 42.153 42.854 40.220 38.756	52.766 51.348 50.971 53.545 54.907 55.675 53.569 53.180 53.654 53.001 55.234 54.964 50.596 49.299 48.308 47.306 49.519 48.252 48.565 47.651 46.595 45.813 49.440 47.150 48.329 46.568 45.729 44.771 45.343 45.469	45.003 45.300 46.419 44.582 43.984 44.992 44.734 46.010 43.277 45.920 45.404 43.806 43.267 43.507 44.208 44.192 45.115 45.551 42.749 42.522 43.365 44.561 42.003 42.750 42.358 45.636 46.284 45.055 46.930 44.280 43.613 43.336 42.184 42.138	16.92 16.63 17.05 21.77 30.11 37.62 42.75 39.25 20.00 20.00 20.00 20.00 16.50 17.63 20.41 21.90 14.84 13.92 20.00	
3318	SER	Н	39.321	47.129	43.860	20.00	

2210	C C D	t-1 A	40 EC 4		44 500		44.404		20.00
3319	SER	HA	40.564		44.582	٠.	44.184		20.00
3320	SER	1HB	40.511		44.313		41.903		20.00
3321	SER	2HB	40.832		45.875		41.397		20.00
3322	SER	HG	38.200		46.170		42.594	*, *	20.00
3323	LEU	N	42.580		46.968		43.678		22.44
3324	LEU	CA	44.021		47.151		43.601		25.58
3325	LEU	Č,	44.758		46.978		44.971	-	29.18
3326	LEU	0	45.885	٠.				. •	
,		_			47.472		45.142	•	34.54
3327	LEU	CB	44.227		48.551		43.032		23.62
3328		CG	43.771		48.613		41.595		22.66
3329	LEU	CD1	44.790		47.909		40.682		23.63
3330		CD2	43.739		50.030		41.191	*	25.59
3331	LEU	H	41.920	1.	47.692		43.869	. ,	20.00
3332	LEU	HA	44.457	-	46.425		42.916		20.00
3333	LEU	1HB	45.272	,	48.860		43.086		20.00
3334	LEU	2HB	43.668		49.264		43.642	•	20.00
3335		HG	42.768	1 -	48.203	•"	41.473		20.00
3336	LEU	1HD1			46.837		40.861		20.00
3337			45.797	•	48.314		40.814	**	
	LEU								20.00
			44.523	1.0	48.057		39:636	. "	20.00
3339			44.730	* .	50.484		41.260	,	20.00
3340			43.082		50.610		41.825	,	20.00
3341	LEU		43.390		50.156		40.166		20.00
3342		Ν	44.047	٠	46.362		45.949		27.88
3343	SER	CA	44.527	-	46:315		47.294	*	28.98
3344	SER	C	45.268		45.026		47.535		29.17
3345	SER	. O	44.928		44.000	* *	46.945		29.28
3346	SER	СВ	43.294		46.473		48.203		29.56
3347	SER	OG	42.391		47.648	· · .	47.851		38.82
3348		Н	43.142		45.971		45.753		20.00
3349	SER	НА	45.282	1	47.090		47.412	•	20.00
3350	SER	1HB	43.680	-,	46.480		49.240		20.00
3351	SER	2HB	42.704	٠.					
3352					45.533		48.129		20.00
4	SER	HG	42.660	,	48.585	1.	47.609		20.00
3353	PRO	N	46.350		45.122	٠,.	48.386		28.99
		CA	47.235		44.009	*	48.606		28.15
3355	PRO	C -	46.665		42.947		49.550		27.34
3356	PRO	0 .	47.292	4.	41.932		49.843		28.43
3357	PRO	CB	48.430		44.664		49.261		29.14
3358	PRO	CG	47.828		45.786		50.081.	2	28.91
3359	PRO	CD [	46.762		46.306		49.144		28.77
3360	PRO	HA	47.507		43.522	0	47.670	vi i	20.00
3361	PRO		49.098		45.079		48.505		20.00
3362	PRO	2HB	49.021		43.976		49.861		20.00
3363		1HG	47.380	* .	45.394		50.989		20.00
3364	PRO	2HG	48.563				50.377	•	
3365				14	46.532				20.00
			47.199 ~		47.044		48.471		20.00
3366	PRO	2HD	45.931		46.753		49.695		20.00
3367	GLU	N	45.457		43.150		49.997		25.89
3368		CA	44.871		41.988		50.605	1	27.19
3369		C	44.368	-	41.012		49.552		26.00
3370	GLU	0	43.942		39.916		49.893		25.89

					·				
3371	GLU	CB	43.911		42.425		51.728		33.55
3372	GLU	CG	42.550		43.115		51.372		41.02
3373	GLU	CD	42.705		44.505		50.654		47.74
3374	GLU	OE1	43.847		45.031		50.685		49.25
3375	GLU	OE2	41.702		45.025		50.090		51.04
							49.758		20.00
3376	GLU	H	44.923		43.947				
3377	GLU	HA	45.628		41.417		51,145		20.00
3378	GLU	1HB	44.457		43.059		52.424		20.00
3379	GLU	2HB	43.665		41.517		52.278		20.00
3380	GLU	1HG	41.959		43,265		52.272		20.00
3381	GLU 1	2HG	41.958		42.466		50.731		20.00
3382	HIS	N	44.471		41.415		48.248		21.92
3383	HIS	CA	44.127		40.631		47.048		18.96
3384	HIS	C	45.367		40.200		46.240		17.83
3385	HIS	Ŏ.	46.450		40.745		46.401		17.07
3386	HIS	CB '	43.211		41.449	•	46.115		20.94
					41.788		46.796	* •	20.70
3387	HIS	CG	41.922						
3388	HIS	ND1	41.464	,	43.035		47.020		22.42
3389	HIS	CD2	40.920		40.898	•	47.210		19.89
3390	'HIS	CE1	40.213	•	42.908		47.536		18.97
3391	HIS	NE2	39.847		41.615		47.661		19.93
3392	HIS	Н	44.691		42.370		48.072		20.00
3393	HIS	HA	43.622		39.722		47.371	**	20.00
3394	HIS	1HB	42.958		40.907		45.199		20.00
3395	HIS	2HB	43.680		42.390		45.807		20.00
3396	HIS	HD1	41.918		43.894		46.893		20.00
3397	HIS	HD2	41.004	•	39.819		47.163		20.00
3398	HIS	HE1	39.593		43.730		47.851		20.00
3399	GLY	N	45.184		39.221		45.302		16.31
3400	GLY	CA	46.302		39.054		44.354		14.07
3401	GLY	C	46.385		40.252	٠.,	43.419		12.35
3402	GLY	Ö	45.650		41.217		43.559		13.46
3403	GLY	H	44.287		38.802		45.246		20.00
	GLY		46.107		38.168		43.769		20.00
3404		1HA				•			20.00
3405	GLY	2HA	47.243		38.952		44.893		11.56
3406	PRO	N	47.313		40.254		42.476		
3407		•	47.457		41.398	-	41.621		13.06
3408	PRO	C.	46.319		41.572		40.614		12.70
3409		0	45.791		40.570	:	40.113		13.68
3410	PRO	CB	48.793		41.186		40.861	1	11.82
3411	PRO	CG	49.194		39.809		41.186		12.42
3412	PRO	CD	48.405		39.318		42.377		11.92
3413	PRO	HA ·	47.521		42.314		42.213		20.00
3414	PRO	1HB	49.530		41.880		41.216		20.00
3415	PRO	· 2HB	48.701		41.370		39.795		20.00
3416	PRO	1HG	48.850		39.180		40.380		20.00
3417	PRO	2HG	50.270		39.657		41.272		20.00
	PRO		49.014		39.338		43.284		20.00
3419	PRO	2HD	48.052		38.297		42.234		20.00
3420	VAL	N:	45.973		42.849		40.334		12.70
3421	VAL	CA	44.929		42.963		39.339		12.91
3422	VAL	C	45.483		42.510		37.953		13.19
U-722	V / \L	_	10.400	•	12.010		JUUŲ		. 5. 10

3423 3424 3425 3426 3427 3428 3429 3431 3432 3433 3433 3433 3433 3444 3445 3445	VVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVVV	1HG2 3HG2 3H C C O C C C H H H H 12 3 H C C O C C C H H H H 12 3 H C C O C C C H H H H 12 3 H C C O C C C H H H H 12 3 H C C C C C C C C C C C C C C C C C C	43.120 45.922 46.576 45.433 44.602 44.881 44.531 43.397 44.176 44.743 44.361 43.709 45.954 43.115 44.479 45.824 44.325 45.410 43.899 43.873 45.479 45.213 46.520 46.535 46.096 46.058 45.692 46.407 44.159 47.120 46.514 45.048 46.421 44.588 44.843	42.728 44.421 44.731 45.279 43.579 42.315 44.757 44.202 44.460 45.801 45.076 45.157 46.341 41.927 42.778 43.158 40.272 39.925 41.713 40.379 39.925 41.413 40.379 39.925 41.413 40.379 42.856 43.856 43.856 45.956 45.956 45.956 46.804 47.202 46.804 47.202 46.804 47.202 46.804 47.202 46.804 47.202 46.804 47.202 46.804 47.403 47		37 620 39 253 38 248 38 707 40 884 39 632 40 258 38 259 37 265 38 259 37 265 38 273 37 38 259 37 38 259 37 38 38 38 38 38 38 38 38 38 38 38 38 38		11.64 13.06 14.25 13.39 20.00
3470 3471	HIS HIS	<u>C</u>	44.257 43.285	 44.454 45.158		28.566 28.830		10.06 9.96
3472	HIS	CB	44.287	41.924	۳.	29.313		8.91
3473 3474	HIS HIS	CG ND1	42.824 42.370	41.948 41.918		28.937 27.656	,	8.64 7.44

			)		101	
3475	HIS	CD2	41.783	41.815	29.793	8.51
3476	HIS	CE1	41.082	41.735	27.708	9.19
3477	HIS	NE2	40.730	41.677	28.984	11.46
3478	HIS	Н	43.749	44.180	31.164	20.00
3479	HIS	HA	45.921	43.222	29.367	20.00
3480	HIS	1HB	44.424	41.327	30.215	20.00
3481	HIS	2HB	44.812	41.427	28.502	20.00
3482	HIS	HD1	42.926	41.960	26.851	20.00
3483	HIS	HD2	41.793	41.781	30.873	20.00
3484	HIS	HE1	40.391	41.585	26.885	20.00
3485	CYS	N	44.901	44.539	27.410	10.31
3486	CYS	CA	44.335	45.149	26.196	8.62
3487	CYS	C	44.351	44.032	25.182	9.41
3488	CYS	Ö	44.133	42.849	25.460	9.11
3489	CYS	CB.	45.153	46.362	25.753	9.94
3490	CYS	SG	47.021	46.150	25.889	10.94
3491	CYS	Η .	45.698	43.931	27.402	20.00
3492	CYS	HA	43.311	45.451	26.378	20.00
3493	CYS	1HB	44.769	47.265	26.233	20.00
3494	CYS	2HB	44.846	46.673	24.780	20.00
3495	CYS	HG	47.564	46.018	24.675	20.00
3496	SER	N	44.663	44.316	23.984	9.15
3497	SER	CA	44.722	43.139	23.130	8.78
3498	SER	C	46.105	42.359	23.239	8.55
3499	SER	Ö	46.103	41.128	23.347	8.15
3500	SER	СВ	44.389	43.699	21.683	7.93
3501	SER	OG	44.662	42.684	20.631	8.26
3502	SER	Н	44.850	45.208	23.621	20.00
3503	SER	HA	43.891	42.482	23.434	20.00
3504	SER	1HB	45.102	44.543	21.547	20.00
3505	SER	2HB	43.440	44.339	21.661	20.00
3506	SER	HG	44.215	41.792	20.609	20.00
3507	ALA	N	47.234	43.166	23.239	8.61
3508		CA	48.571	42.556	23.451	7.66
3509	ALA	C	49.048	42.526	24.949	8.82
3510	ALA	Ŏ.	49.906	41.719	25.386	11.75
3511	ALA	СВ	49.495	43.448	22.703	7.00
3512	ALA	Н	47.101	44.074	22.832	20.00
3513	ALA	HA	48.580	41.555	23.021	20.00
3514	ALA	1HB	49.449	44.478	23.062	20.00
3515	ALA	2HB	49.221	43.481	21.650	20.00
3516	ALA	3HB	50.531	43.112	22.754	20.00
3517	GLY	N	48.451	43.474	25.731	8.66
3518	GLY	CA	48.838	43.523	27.166	8.32
3519	GLY	С	50.020	44.449	27.521	9.76
3520	GLY	0	50.750	44.217	28.493	10.12
3521	GLY	H	47.716	43.998	25.323	20.00
3522	GLY	1HA	49.118	42.517	27.472	20.00
3523	GLY	2HA	47.962	43.705	27.760	20.00
3524	ILE	Ν	50.209	45.476	26.649	11.14
3525	ILE	CA	51.393	46.367	26.588	12.07
3526	ILE	С	51.104	47.859	26.189	10.76

3527 3528 3529 3530 3531 3532 3533 3533 3534 3535 3536 3537 3542 3543 3544 3545 3546 3547 3548 3550 3551 3552 3553 3556 3567 3568 3569 3570 3571	ILILILILILILILILILILILILILICOGOGOGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	1HG2 2HG2	51.483 52.535 52.223 52.925 53.481 49.535 51.746 53.406 51.822 52.130 53.153 53.807 53.265 53.262 50.417 50.161 49.289 49.707 50.328 49.707 47.128 45.607 47.128 45.607 47.128 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47.000 47.139 47		48.766 45.757 45.596 44.361 45.306 45.515 46.417 46.400 46.511 43.636 44.35 43.971 44.318 46.026 45.368 48.135 49.536 50.270 51.067 47.389 49.555 50.422 49.944 50.731 50.827 49.833 49.536 50.223 49.447 51.506 50.223 49.427 51.924 51.		26.906 25.705 24.220 26.249 23.388 25.921 27.621 25.807 23.805 24.044 26.107 27.313 25.748 23.568 23.562 24.720 25.787 26.645 24.405 24.405 24.405 24.405 24.405 24.720 25.787 26.563 28.993 26.013 24.746 24.185 23.356 24.746 24.185 25.769 26.769 27.726 26.769 27.726 27	12.85 8.68 9.17 8.98 7.46 20.00 20.0
3570	ARG	1HH1	41.396		48:399		21.816	20.00
3571 3572			41.590 42.793		49.702 47.113		22.945 21.366	20.00
3573	ARG	2HH2	44.482		47.294	٠	21.669	20.00
3574 3575	SER SER	N . CA	47.594 47.913	- 4	48.689 48.208		28.346 29.693	 10.07 8.26
3576	SER	C	49.065		48.962		30.267	10.19
3577	SER	0	48.901		49.340		31.398	10.34
3578	SER	CB	48.033		46.638		29.836	8.89

3628 CYS C 50.699 52.006 35.192 10.89 3629 CYS O 51.065 52.292 36.335 12.32 3630 CYS CB 50.376 49.689 34.433 12.17		3617 PHE H       47.740       51.934       30.607       20.00         3618 PHE HA       46.395       53.324       32.576       20.00         3619 PHE 1HB       46.680       50.256       32.197       20.00         3620 PHE 2HB       45.487       51.288       31.473       20.00	3612 PHE CD1 43.998 51.687 33.724 11.54 3613 PHE CD2 45.625 50.250 34.736 11.23 3614 PHE CE1 43.198 51.577 34.840 6.44 3615 PHE CE2 44.807 50.110 35.884 9.94	3607       PHE       CA       46.998       52.422       32.513       10.60         3608       PHE       C       47.763       52.282       33.849       10.58         3609       PHE       O       47.468       53.018       34.785       11.24         3610       PHE       CB       46.107       51.169       32.361       8.88	3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.611 55.422 29.584 20.00 3605 THR 3HG2 47.726 55.537 27.806 20.00 3606 PHE N 47.836 52.592 31.360 9.10	3600 THR HA 50.475 54.060 29.842 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3602 THR HG1 49.688 52.952 26.910 20.00	3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73	3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10	3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00	3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00	3582 3583 3584 3585 3586 3587 3588 3590 3591 3593 3594 3595 3599 3600 3601 3602 3603 3604 3605 3606 3616 3616 3617 3618 3618 3621 3622 3623 3624 3628 3629	RRRRRYYYYYYYK RRRRRRRRRRRRRRR HHHHHHHHHH	12HNCCOH12NCCOCOCHHHH1123NCCOCCCCCHH112HHHHHNCCOBBBBCCOHH112NCCOCCCCHH112HHHHNCCOBBBBCCOCCCCHH112HHHHNCCO	48.069 49.068 46.279 50.182 50.896 51.325 50.896 51.649 50.155 49.632 48.658 49.061 47.726 46.998 47.763 47.468 47.740 46.998 47.740 46.998 47.740 46.584 49.631 50.699 51.065	46.443 46.298 46.160 49.229 49.953 51.373 51.879 48.899 49.991 49.439 52.015 53.367 53.501 54.330 53.726 53.774 55.179 51.610 53.223 52.952 55.745 55.422 55.537 52.592 52.422 53.018 51.035 51.687 50.250 51.288 51.934 53.324 50.256 51.288 52.276 49.739 52.006 52.292	30.939 29.561 28.550 29.505 30.405 31.498 29.311 30.989 29.454 29.813 31.196 32.043 28.649 27.401 28.705 28.544 29.842 28.821 26.910 28.772 29.584 27.806 32.513 33.849 34.785 32.876 32.513 33.724 34.736 34.840 35.840 35.919 36.733 36.733 36.793 36.793 36.793 36.793 36.793 36.335	20.00 20.00 10.44 8.60 9.62 11.83 20.00 20.00 11.37 10.02 11.10 9.82 9.53 10.14 8.73 20.00 20.00 20.00 20.00 20.00 20.00 20.00 10.58 11.24 8.88 9.58 11.54 11.23 6.47 20.00 20
3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3591 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3596 THR C 99.111 54.330 32.043 9.82 3596 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR HA 50.061 51.610 28.544 20.00 3600 THR HA 50.475 54.060 29.842 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3602 THR HB 47.638 53.223 28.821 20.00 3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.611 55.422 29.584 20.00 3606 PHE N 47.836 52.592 31.360 9.10 3607 PHE CA 46.998 52.422 32.513 10.60 3608 PHE C 47.763 52.282 33.849 10.58 3611 PHE CG 45.222 51.035 33.635 9.58 3612 PHE CD1 43.998 51.687 33.724 11.54 3613 PHE CG2 44.807 50.110 32.849 6.44 3615 PHE CB 48.680 50.256 32.197 20.00 3617 PHE HB 47.740 51.934 30.607 20.00 3619 PHE C HA 31.998 51.687 33.724 11.54 3619 PHE HB 47.740 51.934 30.607 20.00 3619 PHE D 47.740 51.934 30.607 20.00 3622 PHE HB 46.680 50.256 32.197 20.00 3624 PHE HB 46.680 50.256 36.733 20.00 3625 PHE HZ 42.947 50.692 36.793 34.885 10.02	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3595 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR H 50.061 51.610 28.544 20.00 3601 THR HA 50.475 54.060 29.842 20.00 3603 THR HG1 49.688 52.952 26.910 20.00 3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.611 55.422 39.584 20.00 3606 PHE N 47.836 52.592 31.360 9.10 3607 PHE CA 46.998 52.422 32.513 10.60 3608 PHE C 47.766 55.537 27.806 20.00 3609 PHE O 47.468 53.018 34.785 11.24 3610 PHE CB 46.107 51.169 32.361 8.88 3611 PHE CG 45.222 51.035 33.635 9.58 3612 PHE CD1 43.998 51.687 33.724 11.54 3613 PHE CG2 44.807 50.110 35.884 9.94 3616 PHE CZ 43.575 50.768 35.919 6.47 3617 PHE H 47.740 51.934 30.607 20.00 3620 PHE 1 HB 46.680 50.256 32.197 20.00 36361 PHE HA 46.395 53.324 32.576 20.00 36361 PHE HA 46.395 53.324 32.576 20.00 3640 PHE D 47.468 50.018 35.919 6.47	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY D 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR H 50.061 51.610 28.544 20.00 3600 THR HA 47.638 53.223 28.821 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3602 THR HG1 49.688 52.952 26.910 20.00 3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.726 55.537 27.806 20.00 3605 THR 3HG2 47.726 55.537 27.806 20.00 3606 PHE N 47.836 52.592 31.360 9.10 3607 PHE CA 46.998 52.422 32.513 10.60 3608 PHE C 47.763 52.282 33.849 10.58 3610 PHE CB 46.107 51.169 32.361 8.88 3611 PHE CG 45.222 51.035 33.635 9.58 3612 PHE CD1 43.998 51.687 33.724 11.54 3613 PHE CD2 45.625 50.250 34.736 11.23 3614 PHE CE1 43.198 51.577 34.840 6.44 3615 PHE CE2 44.807 50.110 35.884 9.94	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR H 50.061 51.610 28.544 20.00 3600 THR HA 50.475 54.060 29.842 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3602 THR HG1 49.090 55.745 28.772 20.00 3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.611 55.422 39.584 20.00 3606 PHE N 47.836 52.592 31.360 9.10 3607 PHE CA 46.998 52.422 32.513 10.60 3608 PHE C 47.763 52.282 33.849 10.58 3609 PHE O 47.468 53.018 34.785 11.24 3610 PHE CB 46.107 51.169 32.361 8.88	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR H 50.061 51.610 28.544 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3603 THR 1HG2 49.090 55.745 28.772 20.00 3604 THR 2HG2 47.611 55.422 29.584 20.00	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR CG 48.846 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73 3599 THR H 50.061 51.610 28.544 20.00 3600 THR HA 50.475 54.060 29.842 20.00 3601 THR HB 47.638 53.223 28.821 20.00 3602 THR HG1 49.688 52.952 26.910 20.00	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196 11.10 3595 THR O 49.111 54.330 32.043 9.82 3596 THR CB 48.658 53.726 28.649 9.53 3597 THR OG1 49.394 53.774 27.401 10.14 3598 THR CG2 48.212 55.179 28.705 8.73	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3591 GLY 2HA 51.649 49.439 30.989 20.00 3592 THR N 50.155 52.015 29.454 11.37 3593 THR CA 49.632 53.367 29.813 10.02 3594 THR C 48.846 53.501 31.196	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62 3588 GLY O 51.139 51.879 31.498 11.83 3589 GLY H 50.224 48.899 28.558 20.00 3590 GLY 1HA 52.103 49.991 29.311 20.00 3580 GLY 1HA 52.103 49.991 29.311	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00 3584 SER HG 46.279 46.160 28.550 20.00 3585 GLY N 50.182 49.229 29.505 10.44 3586 GLY CA 51.325 49.953 30.078 8.60 3587 GLY C 50.896 51.373 30.405 9.62	3581 SER HA 47.089 48.557 30.322 20.00 3582 SER 1HB 48.069 46.443 30.939 20.00 3583 SER 2HB 49.068 46.298 29.561 20.00								1

3631 3632 3633 3634 3635 3636 3637 3638 3640 3641 3642 3643 3644 3645 3650 3651 3652 3653 3654 3655 3656 3667 3668 3667 3668 3670 3671 3672 3673 3674	CCCCCLLLLLLLLLLLLLLLLLLLLAAAAAAAAAAAAA		49.243 48.709 49.055 51.208 50.793 48.768 51.226 52.185 51.674 52.319 52.734 53.305 54.937 55.470 54.636 54.937 55.470 54.680 55.757 54.680 55.757 54.680 55.757 56.511 57		48.345 50.767 50.755 49.519 49.726 48.009 52.596 53.673 54.833 55.313 55.313 54.327 56.096 52.331 54.585 53.374 56.567 55.435 56.419 56.419 56.849 57.733 57		34.666 32.964 35.790 35.117 33.426 33.459 34.112 34.273 35.143 36.051 32.895 31.977 34.036 33.223 34.825 32.286 32.332 33.544 35.013 34.725 35.284 36.768 37.671 34.272 33.918 35.359 34.616 37.028 38.415 39.379 40.421 38.350 39.753 40.031 36.245		15.41 20.00 20.00 20.00 20.00 20.00 10.01 10.37 10.12 9.30 9.12 11.23 11.32 11.96 20.00 20	
3671	ASP	CG	47.508		52.440		39.753		13.48	
100										
3675	ASP	HA	47.781		55.068		38.858	,	20.00	
3676	ASP	1HB	48.599		52.290		37.857		20.00	
3677	ASP	2HB	47.011	-	53.008		37.763 <sup></sup>		20.00	,
3678		N	50.869_		.53.897_	•	38.951_		10.97_	
3679		CA	52.060		53.792		39.783		11.02	
3680	THR	C	52.733		55.095		39.943		11.79	
3681	THR	Ö	53.143		55.434		41.062		11.99	
3682	THR	CB	53.122		52.827	•	39.159		11.07	

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3787	ASP	1HB	50.143	57.792	46.991	20.00
3788	ASP	2HB	49.523	58.841	45.757	20.00
3789	LYS	N	52.983	58.760	48.131	22.67
3790	LYS	CA	53.674	58.145	49.290	26.79
3791	LYS	C	54.931	58.920	49.795	27.91
3792	LYS	0	55.952	58.446	50.319	27.25
3793	LYS	СВ	53.901	56.634	49.079	31.63
3794	LYS	CG	55.221	56.174	48.432	36.21
3795	LYS	CD	55.564	54.732	48.920	41.15
3796	LYS	CE	54.502	53.672	48.501	45.83
3797	LYS	NZ	54.382	52.509	49.411	48.52
3798	LYS	Н	53.168	58.421	47.202	20.00
3799	LYS	HA .	52.971	58.231	50.127	20.00
3800	LYS	1HB	53.037	56.222	48.565	20.00
3801	LYS	2HB	53.878	56.200	50.076	20.00
3802	LYS	1HG	56.072	56.800	48.691	20.00
					· ·	
3803	LYS	2HG	55.142	56.208	47.346	20.00
3804	LYS	1HD	55.694	54.725	50.001	20.00
3805	LYS	2HD	56.528	54.416	48.507	20.00
3806	LYS	1HE	54.738	53.304	47.496	20.00
3807	LYS	2HE	53.519	54.136	48.398	20.00
3808	LYS	1HZ	54.131	52.810	50.372	20.00
3809	LYS	2HZ	55.273	51.969	49.417	20.00
3810	LYS	3HZ	53.633	51.882	49.045	20.00
3811	ARG:	N ·	54.783	60.192	49.561	29.18
3812	ARG	CA	55.920	61.072	49.744	30.25
3813	ARG	С	55.412	62.476	49.597	28.51
3814	ARG	0	55.983	63.400	50.172	29.83
3815	ARG	СB	57.138	60.726	48.833	35.96
3816	ARG	CG	56.825	60.702	47.353	35.60
3817	ARG	CD	57.971	60.140	46.489	39.72
3818	ARG	NE	58.346	58.740	46.746	44.72
3819	ARG	CZ	59.252	58.052	45.985	49.49
3820	ARG	NH1	59.736	58.571	44.823	46.55
3821	ARG	NH2	59.631	56.831	46.444	54.27
3822	ARG		53.921	60.398	49.096	20.00
3823	ARG	HA	56.222	60.959	50.792	20.00
3824	ARG	1HB	57.517	59.753	49.136	20.00
3825	ARG	2HB	57.954	61.419	49.015	20.00
3826	ARG	1HG	56.572	61.699	46.997	20.00
3827	ARG	2HG	55.957	60.080	47.195	20.00
3828	ARG	1HD	58.897	60.700	46.658	20.00
3829	ARG	2HD	57.668	60.138	45.441	20.00
3830	ARG	HE	57.974	58.277	47.547	20.00
3831	ARG	1HH1	60.317	57.968	44.264	20.00
3832	ARG		59.524	59.501	44.534	20.00
3833	ARG	,	60.241	56.297	45.852	20.00
			59.320	56.463	47.308	20.00
3835	LYS	N	54.299	62.635	48.843	24.03
3836	LYS	CA	53.709	63.975	48.683	22.66
3837	LYS	C	54.836	65.076	48.313	22.19
3838	LYS	Ö	54.746	66.296	48.531	22.54
5050	_10	•	UT.17U	55.255	70.00 i	£2.07

3839	LYS	СВ	52.828	64.382	49.905	23.45
3840		CG	51.703	63.412	50.390	20.10
3841	LYS	CD	50.532	63.202	49.448	20.71
3842	LYS	CE	49.505	62.232	50.059	17.94
3843		NZ	48.450	61,958	49.071	27.72
3844	LYS	Н	53.968	61.880	48.276	20.00
3845	LYS	HA	53.071	63.873	47.806	20.00
3846		1HB	52.363	65.342		
					49.684	20.00
3847	LYS	2HB	53.494	64.560	50.747	20.00
3848	LYS	1HG	51.304	63.828	51.315	20.00
3849	LYS	2HG	52.136	62.454	50.672	20.00
3850	LYS	1HD	50.884			
				62.812	48.489	20.00
3851	LYS	2HD	50.052	64.161	49.235	20.00
3852	LYS	1HE	49.049	62.670	50.950	20.00
3853	LYS	2HE	49.952	61.283	50.363	20.00
3854	LYS	1HZ	48.810	61.313	48.331	20.00
					*/	
3855	LYS	2HZ	48.109	62.817	48.578	20.00
3856	LYS	3HZ	47.647	61.409	49.432	20.00
3857	ASP	Ν	55.889	64.482	47.680	21.48
3858	ASP	CA	56.806	65.251	46.882	22.85
3859	ASP	C				
			56.543	64.944	45.345	21.41
3860	ASP	0	57.198	64.103	44.713	21.48
3861	ASP	CB	58.216	64.968	47.416	25.58
3862	ASP	CG	59.244	65.767	46.566	29.64
3863	ASP	OD1	58.862	66.819	45.964	31.77
3864	ASP	OD2	60.408	65.344	46.549	30.41
3865	ASP	Н	56.060	63.508	47.768	20.00
3866	ASP	HA	56.609	66.305	47.057	20.00
3867	ASP	1HB	58.465	63.91.8	47.556	20.00
3868	ASP					
		2HB	58.277	65.369	48.429	20.00
3869	PRO	Ν	55.539	65.638	44.744	20.58
3870	PRO	CA	55.308	65.416	43.295	22.04
3871	PRO	С	56.569	65.307	42.347	24.22
3872	PRO	Ŏ.	56.572	64.547	41.372	22.77
		_		•		
3873	PRO	СВ	54.441	66.653	42.852	23.82
3874	PRO	CG	53.958	67.350	44.156	22.76
3875	PRO	CD	54.861	66.824	45.284	22.17
3876	PRO	HA	54.784	64.489	43.157	20.00
3877	PRO	1HB	53.643	66.360	42.176	and the second s
						20.00
3878	PRO	2HB	55.022	67.367	42.285	20.00
3879	PRO	1HG	53.966	68.430	44.056	20.00
3880	PRO	2HG	52.945	67.022	44.344	20.00
3881	PRO	1HD	54.292	66.563	46.162	20.00
3882	PRO	2HD	55.589	67.586	45.557	20.00
3883	SER	N	57.644	66.017	42.810	26.49
3884	SER	CA	58.744	66.328	41.960	29.21
3885	SER ·		59.732	65.218	41.913	28.16
3886	SER	Ö	60.301			
				64.849	40.890	30.98
3887	SER	СВ	59.304	67.616	42.594	32.36
3888	SER	OG	58.637	68.913	42.285	37.57
3889	SER	Н	57.630	66.394	43.738	20.00
3890	SER	HA .	58.429	66.385	40.908	20.00
2000	JLIN	11/3,	JU.720	00.303	70.300	20.00

					400	
3891	SER	1HB	60.351	67.694	42.232	20.00
3892	SER	2HB	59.521	67.454	43.682	20.00
3893	SER	HG	57.670	69.067	42.060	20.00
3894	SER	N	59.925	64.613	43.043	25.94
3895	SER	CA	60.809	63.466	42.969	23.97
3896						
	SER	C	60.407	62.308	41.968	23.07
3897	SER	0	61.251	61.419	41.868	23.41
3898	SER	CB	60.695	62.845	44.395	24.53
3899	SER	OG	59.325	62.461	44.856	26.18
3900	SER	Н	59.459	64.900	43.888	20.00
3901	SER	HA	61.810	63.811	42.723	20.00
3902	SER	1HB	61.331	63.448	45.113	20.00
3903	SER	2HB	61.358	61.949	44.388	20.00
3904	SER	HG	58.514	63.013	44.586	20.00
3905	VAL	Ν	.59.176	62.231	41.306	22.45
3906	VAL	· CA	58.834	61.027	40.461	21.20
3907	VAL	С	59.182	61.249	38.947	21.26
3908	VAL	0	58.675	62.163	38.320	22.49
3909	VAL	СВ	57.391	60.434	40.620	22.58
3910	VAL	CG1	56.451	60.386	39.363	20.76
3911	VAL	CG2	56.695	61.029	41.828	17.17
3912	VAL	Н	58.588	63.040	41.291	20.00
3913	VAL	HA	59.500	60.233	40.807	20.00
3914	VAL	HB	57.535	59.380	40.863	20.00
3915	VAL	1HG1		59.787	38.559	20.00
3916	VAL	2HG1		61.381	38.973	20.00
3917	VAL		55.488	59.919		
					39.577	20.00
3918	VAL		56.539	62.101	41.733	20.00
3919	VAL		57.252	60.803	42.729	20.00
3920	VAL		55.729	60.556	41.956	20.00
3921	ASP	N	60.034	60.370	38.393	19.92
3922	ASP	CA	60.422	60.486	37.014	19.98
	ASP	C	59.593	59.489	36.190	17.61
3924		0	59.810	58.291	36.270	18.51
	ASP	СВ	61.968	60.298	36.922	21.39
3926	ASP	CG	62.632	60.452	35.517	25.31
3927	ASP	OD1	62.006	60.671	34.458	23.58
3928	ASP	OD2	63.846	60.336	35.498	32.08
3929	ASP	Н	60.462	59.701	38.992	20.00
3930	ASP	HA	60.181	61.487	36.644	20.00
3931	ASP	1HB	62.267	59.369	37.393	20.00
3932	ASP	2HB.	62.408	61.078	37.534	20.00
3933	ILE	Ν -	58.641	60.060	35.429	18.00
3934	ILE	CA	57.715	59.212	34.684	16.17
3935	ILE	C	58.457	58.121	33.908	16.20
3936	ILE	Ö	58.158	56.962	33.989	16.71
3937	ILE	СВ	56.682	60.059	33.831	18.69
3938	ILE	CG1	55.862	61.009	34.788	20.07
3939	ILE	CG2	55.633	59.144	33.082	16.13
3940	ILE	CD1	54.494	61.498	34.162	22.35
3941	ILE	Н	58.643	61.059	35.410	20.00
3942	ILE	HA	57.154	58.675	35.450	20.00
JJ42			57.154	30.073	33,430	20.00

3966 3967 3968 3969 3970 3971 3972 3973 3974 3975 3976 3977 3978 3979 3980	LYS LYS LYS LYS LYS LYS LYS LYS LYS LYS	2HG2 3HD1 3H N C C O C C C N H H 1H H H H H H H H H H H H H H H H	55.008 56.178 54.982 53.818 54.710 53.924 59.488 60.319 61.173 61.228 61.147 60.600 61.795 62.746 62.754 59.666 59.593 61.394 62.114 60.123 59.834 61.613 62.379 63.379 63.370 62.462 61.799 63.031 63.326 61.763 62.721 63.685 64.794		60 649 61 858 60 471 58 573 58 446 59 699 60 694 62 104 58 562 57 703 56 727 55 568 58 457 59 764 60 577 61 132 62 577 59 540 57 072 57 754 58 664 60 303 59 586 61 325 59 879 60 714 62 973 63 140 62 973 63 140 65 7 189 57 072 57 072	33.091 35.081 35.081 35.709 33.774 32.443 32.406 33.876 33.280 34.865 33.158 32.288 33.130 32.799 31.166 30.558 29.985 31.087 31.255 33.256 31.780 30.371 31.609 31.374 29.806 29.225 29.384 30.916 32.058 31.456 30.419 32.058 31.456 30.419 32.070 34.225 35.671 35.780 36.7420 37.285 37.807 36.734		20.00 20.00 20.00 20.00 20.00 20.00 20.00 20.00 15.61 16.12 15.62 15.12 18.84 31.31 37.73 43.40 47.06 20.00
3972	LYS	2HZ	63.031		63.140	 30.419		20.00
3974	LYS	Ν	61.763		57.189	34.225		16.19
3977	LYS	0	61.750		53.857	35.780		13.38
3979 3980	LYS	CG CD	63.685		56.179	37.285	. •	26.41
3981 3982 3983	LYS LYS LYS	CE NZ H	65.626 67.041 61.705		57.849 58.078 58.168	36.734 37.149 34.443	,	37.01 41.44 20.00
3984 3985	LYS LYS	HA 1HB	63.127 61.887	·	55.786 57.362	34.726 37.030		20.00
3986 3987 3988	LYS LYS LYS	2HB 1HG 2HG	63.231 64.127 63.149		57.909 55.380 55.685	36.056 36.694 38.097		20.00 20.00 20.00
3989 3990	LYS LYS	1HD 2HD	65.457 64.355		56.465 57.817	 38.425 38.475_	-	20.00 20.00
3991 3992 3993	LYS LYS LYS	1HE 2HE 1HZ	65.173 65.625 67.519		58.818 57.311 57.178	36.541 35.781 37.364		20.00 20.00 20.00
3994	LYS	2HZ	67.048		58.692	37.990	•	20.00

3995 3996	LYS VAL	3HZ N	67.545 59.985	58.575 55.323	36.382 35.959	20.00 15.38
3997	VAL	CA	59.007	54.265	36.251	13.40
3998	VAL	C	58.622	53.424	35.039	12.42
3999	VAL	ŏ	58.570	52.215	35.094	12.19
4000	VAL	СВ	57.954	54.563	37.383	17.62
4001	VAL	CG1	56.568	53.984	37.220	13.70
4002	VAL	CG2	57.966	55.998	37.910	14.26
4003	VAL	Н	59.750	56.303	35.924	20.00
4004	VAL	HA	59.627	53.531	36.757	20.00
4005	VAL	HB	58.323	54.004	38.246	20.00
4006	VAĽ	1HG1	56.577	52.930	36.941	20.00
4007	VAL		56.042	54.545	36.459	20.00
4008	VAL		55.990	54.074	38.141	20.00
4009	VAL		57.755	56.693	37.096	20.00
4010	VAL		58.937	56.269	38.321	20.00
4011	VAL		57.210	56.147	38.681	20.00
4012	LEU	N	58.433	54.033	33.899	11.53
	LEU	CA	58.144	53.234	32.728	12.42
4014	LEU	С	59.278	52.236	32.445	11.91
4015	LEU	0	59.058	51.051 54.457	32.366	10.65 12.94
4016 4017	LEU LEU	CB CG	57.743 57.326	54.157 53.410	31.624 30.304	14.51
4017	LEU	CD1	57.086	54.408	29.182	14.86
4019	LEU	CD2	56.160	52.386	30.503	12.64
4020	LEU	H	58.437	55.031	33.911	20.00
4021	LEU	HA	57.258	52.666	32.990	20.00
4022	LEU	1HB	58.593	54.808	31.418	20.00
4023	LEU	2HB	56.941	54.822	31.950	20.00
4024	LEU	HG	58.192	52.818	30.000	20.00
4025	ĿEU	1HD1	57.963	55.014	29.049	20.00
4026	LEU	2HD1	56.240	55.060	29.390	20.00
4027	LEU	3HD1	56.896	53.878	28.248	20.00
4028	LEU	1HD2	55.276	52.909	30.864	20.00
4029	LEU		56.404	51.623	31.245	20.00
4030	LEU		55.893	51.869	29.585	20.00
4031	LEU	N	60.504	52.731	32.420	14.10
4032	LEU	CA	61.690	51,872	32.395	13.57
4033	LEU.	С	61.761	50.752	33.479	11.98
4034	LEU	O CB	62.002	49.612	33.082	10.15
4035	LEU	CB	62.918	52.796 52.177	32.471	15.92
4036 4037	LEU LEU	CG CD1	63.686 64.024	53.177 54.658	31.132 31.092	18.96 18.76
4037	LEU	CD2	63.078	52.694	29.832	20.15
4039	LEU	H	60.599	53.726	32.457	20.00
4040	LEU	HA	61.655	51.323	31.457	20.00
4041	LEU	1HB	63.669	52.330	33.103	20.00
4042		2HB	62.658	53.693	_33.033_	20.00
4043		HG	64.658	52.680	31.172	20.00
4044	LEU		64.621	54.977	31.950	20.00
4045	LEU	2HD1	63.112	55.255	31.098	20.00
4046	LEU	3HD1	64.604	54.927	30.204	20.00

4047 4048 4049 4050 4051 4053 4054 4055 4056 4057 4058 4063 4064 4065 4066 4067 4068 4067 4071 4072 4073 4076 4077 4076 4077 4078 4077 4078 4080 4081 4082 4083 4084 4085 4086 4087 4088 4089 4080 4081 4081 4082 4083 4084 4085 4086 4087 4088 4089 4089 4080 4081 4081 4082 4083 4084 4085 4086 4087 4088 4089 4089 4089 4089 4089 4089 4089	UUPPPPPPPPPPPPPTTTTTTTTTTTTTTTTTTGGGGGGGG	2HD2 3H C C C C C O C H H 1 2 N C C O C C S C H H 1 2 1 2 1 2 1 N C C O C C C N C N N N C C O C C C N C N	62.022 63.159 63.545 61.560 61.390 60.231 61.278 62.301 61.278 62.301 61.278 62.301 61.278 62.301 60.255 61.732 58.001 58.462 59.093 54.062 59.129 57.666 55.734 54.045 59.282 59.642 60.637 60.925 59.943 59.943 59.943 59.943 59.943 59.945 59	52.950 51.613 53.169 51.032 49.905 48.923 47.695 50.344 49.288 49.262 48.379 51.996 49.314 50.372 51.315 49.698 50.357 51.043 50.456 48.053 49.008 50.456 48.842 51.743 50.220 51.565 48.214 47.419 46.252 45.526 48.311 49.556 50.017 50.296 51.031 49.731 49.173	29.855 29.727 28.966 34.837 35.801 35.403 35.475 37.304 38.082 38.000 38.689 35.102 35.656 37.670 37.466 34.944 34.411 33.363 33.408 33.836 34.916 34.915 35.256 33.008 35.537 35.576 36.278 36	20.00 20.00 13.06 12.70 11.44 11.16 15.05 23.08 22.88 24.92 20.00
4087	ARG	CZ ·	58.839	50.296	26.192	19.36
4089	ARG	NH2	59.579	49.731	25.327	15.47
4091	ARG ARG	H HA	59.531 58.706	49.173 46.936	32.573 30.961	20.00 20.00
4092 4093	ARG ARG	1HB 2HB	59.788 60.999	47.723 48.599	29.092 29.991	20.00 20.00
4094 4095	ARG	1HG	59.729	50.392	30.254	20.00
4096	ARG ARG	2HG 1HD	58.294 57.908	49.528 50.943	30.682 28.745	20.00 20.00
4097 4098	ARG	2HD HE	57.606 60.107	49.239 49.639	28.396 27.637	20.00 20.00

4151	ARG	CD	53.173	42.198	29.472	8.35
4152	ARG	NE	52.329	41.703	28.413	8.45
4153	ARG	CZ	52.240	40.450	27.919	8.44
4154	ARG ARG	NH1 NH2	52.939	39.527	28.425	8.74
4155 4156	ARG	H	51.413 57.860	40.156 44.166	26.940 32.007	9.54 20.00
4157	ARG	П НА	57.284	41.962	32.007 30.171	20.00
4158	ARG	1HB	55.116	44.056	30.542	20.00
4159	ARG	2HB	55.154	42.528	31.341	20.00
4160	ARG	1HG	55.200	41.356	29.029	20.00
4161	ARG	2HG	54.899	42.903	28.332	20.00
4162	ARG	1HD	52.863	43.247	29.522	20.00
4163	ARG	2HD	52.834	41.843	30.446	20.00
4164	ARG	HE	51.722	42.406	28.035	20.00
4165	ARG	1HH1	52.806	38.586	28.112	20.00
4166	ARG	2HH1	53.602	39.812	29.109	20.00
4167	ARG		51.312	39.214	26.626	20.00
4168	ARG		50.878	40.883	26.495	20.00
4169	MET	N	57.716	43.404	28.075	11.10
4170	MET	CA	58.186	44.260	26.962	10.52
4171	MET	C	57.024	45.223	26.500	12.21
4172	MET	0	55.843	44.858	26.471	11.69
4173	MET	CB	58.606	43.362	25.796	12.40
4174	MET	CG	57.434	42.486	25.214	13.12
4175	MET	SD CE	57.895	41.619	23.717	12.49
4176 4177	MET MET	H	59.286 57.157	40.641 42.590	24.231	14.52
4178	MET	HA	59.033	44.839	27.946 27.332	20.00
4179	MET	1HB	59.421	42.716	26.125	20.00
4180	MET	2HB	59.017	43.981	24.995	20.00
4181	MET	1HG	56.537	43.061	24.993	20.00
4182	MET	2HG	57.153	41.751	25.963	20.00
4183	MET	1HE	58.999	40.071	25.111	20.00
4184	MET	2HE	60.128	41.283	24.473	20.00
4185	MET	3HE	59.569	39.950	23.436	20.00
4186	GLY	N	57.448	46.438	26.060	11.82
4187	GLY	CA	56.696	47.240	25.071 <sup>-</sup>	10.69
4188	GLY	C	55.505	47.941	25.724	11.80
4189	GLY	0	54.605	48.408	25.044	11.81
4190	GLY	H	58.395	46.587	26.326	20.00
4191	GLY	1HA	56.379	46.563	24.277	20.00
4192	GLY	2HA	57.369	47.983	24.659	20.00
4193 4194	LEU	N	55.644	48.052	27.083	11.43
4194	LEU LEU	CA C	54.762 54.798	48.854 50.309	27.881 27.386	11.39 12.20
4196	LEU	0	55.819	51.000	27.369	12.24
4197	LEU	СВ	55.125	48.664	29.353	10.83
4198		CG	55.153	47.177	_29.826	9.47
4199	LEU	CD1	53.979	46.390	29.211	9.16
4200	LEU	CD2	55.106	46.987	31.376	10.38
4201	LEU	Н	56.255	47.429	27.563	20.00
4202	LEU	HA	53.765	48.452	27.684	20.00

4203	LEU	1HB	54.363	49.187		29.928		20.00
4204	LEU-	2HB	56.038	49.177		29.650		20.00
4205	LEU	HG	56.074	46.731		29.458		20.00
4206	LEU	1HD1		46.270	•	28.145		20.00
								20.00
4207	LEU		53.059	46.958		29.330		
4208	LEU		53.845	45.402		29.632		20.00
4209	LEU		54.239	47.489		31.812		20.00
4210	LEU		56.013	47.373		31.822		20.00
4211	LEU	3HD2	55.039	45.931		31.632		20.00
4212	ILE	N	53.611	50.747		26.925		10.54
4213	ILE	CA	53.352	51.904		26.078		11.17
4214	ILE	C	53.942	51.707		24.614		13.75
4215	ILE	0	55.152	51.525		24.341		13.91
4216	ILE <	CB	53:775	53.199		26.797		11.11
4217	ILE	CG1	53.323	53.232		28.236		9.13
4218	ILE	CG2	53.534	54.509		25.973		14.38
4219	ILE	CD1	53.085	54.633		28.731		8.65
4220	ILE	H :	52.857	50.110		27.076		20.00
4221	ILE	HA	52.300	51.849		25.895		20.00
4222	ILE	HB	54.854	53.154		26.854		20.00
4223	ILE	1HG1		52.753		28.875		20.00
						28.395		20.00
4224	ILE		52.434	52.641				
4225	ILE		52.489	54.771		26.013	,	20.00
4226	ILE		53.837	54.423		24.931		20.00
4227	ILE		54.073	55.352		26.405		20.00
4228	ILE		52.339	55.011		28.055		20.00
4229	ILE	2HD1		55.277	,	28.718		20.00
4230	ILE	3HD1	52.642	54.642		29.720		20.00
4231	GLN	N	52.962	51.794		23.698		13.18
4232	GLN	CA	53.199	51.506		22.275		12.31
4233	GLN	С	53.430	52.716		21.367		14.51
4234	GLN	0	53.955	52.560		20.262		14.93
4235	GLN	CB	52.144	50.560		21.736		12.10
4236	GLN	CG	52.439	49.160		22.274		13.08
4237	GLN	CD	53.546	48.547		21.425		14.33
4238	GLN	OE1	53.483	48.340		20.241		16.45
4239	GLN	NE2	54.621	48.324		22.095		11.97
4240	GLN	Н	52.069	52.116		24.025		20.00
4241	GLN	HA	54.053	50.882		22.198		20.00
4242		1HB	52.148	50.567		20.649		20.00
4243	GLN	2HB	51.163	50.906		22.068		20.00
4244	GLN	1HG	51.565	48.532		22.129		20.00
4245	GLN	2HG	52.700	49.128		23.334		20.00
4246	GLN		55.255	47.877		21.463		20.00
4247	GLN		54.687	48.534		23.072		20.00
4248	THR	N	52.999			21.847		14.47
•		CA		53.889 55.112		21.160		13.82
	THR		53.293					
	THR		53.795	 56.277		22.022		15.31
4251	THR	0	53.504	56.378		23.202		15.21
4252	THR	CB	52.018	55.721		20.549		12.44
4253	THR	OG1	51.132	56.387		21.467		13.84
4254	THR	CG2	51.116	54.688		19.900		12.80

4255 4256 4257 4258 4259 4261 4263 4264 4265 4266 4267 4268 4267 4273 4274 4275 4276 4277 4278 4279 4281 4282 4283 4284 4285 4286 4287 4286 4287 4288 4289 4291 4292 4293 4294 4295 4296 4297 4298 4298 4298 4298 4298 4298 4298 4298	THHHHHHAAAAAAAAAAAAAAAAAAAGGGGGGGGGGGGG	H H H H 1 2 3 N C C O C H H 1 2 3 N C C O C C O O H H 1 2 N C C O C C O N H H 1 2 1 H C C O C C C O N H H 1 2 1 1 2 1 2 N C C O C C C O N H H 1 2 1 1 2 N C C O C C C O N H H 1 2 1 1 2 N C C O C C C O N H H 1 2 1 1 2 N C C O C C C O N H H 1 2 1 1 2 N C C O C C O N H H 1 2 1 1 2 N C C O C C O N H H 1 2 1 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C O N H H 1 2 N C C O C C C O N H H 1 1 2 N C C O C C C O N H H 1 1 2 N C C O C C C O N H H 1 1 2 N C C O C C C O N H H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C O N H 1 1 2 N C C O C C C C O N H 1 1 2	51.562 50.153 54.477 54.775 53.591 53.789 55.595 54.845 55.020 56.466 55.932 51.253 50.459 49.950 50.285 49.950 50.285 49.622 48.176 50.796 50.509 50.796 50.509 50.357 51.372 50.983 50.268 49.003 49.169 50.245 48.120 50.220 49.410 50.120 51.199 48.722 48.157 48.192		53.881 54.925 56.216 57.060 53.814 54.409 55.136 57.213 58.515 59.401 60.142 59.324 56.937 58.310 59.526 59.361 60.117 59.397 60.007 60.449 61.573 62.439 61.573 62.439 61.573 62.439 61.573 58.059 57.350 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.732 57.853 57.686 57.687 57.686 57.687 57.686 57.687 57.962 57.962 57.962 57.962 57.962 57.962 57.962 57.962 57.962	22.722 20.382 19.551 22.111 20.510 18.964 19.640 21.304 21.304 22.373 23.290 20.420 22.799 19.991 20.425 23.628 24.564 21.279 21.733 22.425 23.628 24.564 21.279 21.733 22.570 21.255 20.940 22.821 24.885 27.011 24.870 24.131 23.674 23.675 24.675 25.675 26.675 26.675 27		20.00 20.00
							,	
4302	LEU	C	53.022		59.639	 27.103		14.41
4304	LEU	0	53.197		59.793	28.299		15.65
4305	LEU	CB	55.040	٠.	58.373	25.963		12.86
4305	LEU	CG	55.940		58.940	27.044		13.02

4307 4308 4309 4310 4311 4312 4313	LEU LEU LEU LEU LEU	CD1 CD2 H HA 1HB 2HB HG	57.423 55.803 52.853 53.697 55.109 55.368 55.643		59.114 58.093 57.713 57.475 58.990 57.378 59.956		26.568 28.344 24.542 27.137 25.075 25.665 27.299 25.718		12.87 12.32 20.00 20.00 20.00 20.00 20.00 20.00
4314 4315 4316	LEU LEU	1HD1 2HD1 3HD1	57.503 57.826 58.066		59.794 58.155 59.498		26.246 27:365		20.00 20.00
4317 4318 4319	LEU LEU	1HD2 2HD2 3HD2	55.984 54.823 56.521	!	57.030 58.181 58.432		28.190 28.813 29.090		20.00 20.00 20.00
4320 4321	ARG ARG	N CA	52.940 52.516	(	30.581 31.919		26.220 26.552		13.43 13.38
4322 4323 4324	ARG ARG ARG	C O CB	51.245 51.086 52.251	(	61.967 62.719 62.603		27.425 28.402 25.232		14.16 14.64 13.46
4325 4326	ARG ARG	CG CD	51.762 51.576	(	64.060 64.868	•	25.435 24.119		13.29 16.52
4327 4328 4329	ARG ARG ARG	NE CZ NH1	51.522 50.397 49.216	(	66.299 67.002 66.446		<ul><li>24.432</li><li>24.409</li><li>24.187</li></ul>		20.17 19.49 20.88
4330 4331	ARG ARG	NH2 H	50.463 53.137	(	68.265 60.365		24.567 25.265 27.095	,	21.32 20.00 20.00
4332 4333 4334	ARG ARG ARG	HA 1HB 2HB	53.333 51.506 53.170	(	62.403 62.063 62.609		24.656 24.642		20.00
4335 4336 4337	ARG ARG ARG	1HG 2HG 1HD	52.524 50.846 50.719	(	64.576 64.108 64.529	•00	26.019 26.022 23.542	٠,	20.00 20.00 20.00
4338 4339	ARG ARG	2HD HE	52.437 52.358	(	64.758 66.829		23.467 24.622		20.00 20.00
4340 4341 4342	ARG ARG ARG		48.378 49.238 49.696	(	66.959 65.453 68.881		<ul><li>24.120</li><li>24.060</li><li>24.571</li></ul>		20.00 20.00 20.00
4343 4344 4345	ARG PHE PHE	2HH2 N CA	51.389 50.335 49.001	(	68.673 61.083 61.017		24.644 26.968 27.582		20.00 13.41 13.00
4346 4347	PHE PHE	CO	49.112 48.608	(	60.777 61.592		29.110 29.880		12.42 14.84
4348 4349 4350	PHE PHE PHE	CB CG CD1	48.098 46.828 45.801	(	60.098 60.022 60.881		26.822 27.581 27.276		12.87 13.08 15.82
4351 4352	PHE PHE	CD2 CE1	46.705 44.667		59.168 60.910		28.680 28.097		13.48 14.88
4353 4354 4355	PHE PHE PHE	CE2 CZ H	45.574 44.540 50.562	0	59.191 60.071 60.635		29.498 29.199 26.109		14.71 13.48 20.00
4356 4357 4358	PHE PHE PHE	HA 1HB 2HB	48.585 48.536 47.915	:	62.029 59.108 60.462		27.497 26.759 25.813		20.00 20.00 20.00

		*				
4388 4389 4390 4391 4392 4393 4394 4395 4396 4397 4398 4399 4400 4401	LEU	N CA C O CB CG	45 882 47 512 43 859 45 506 43 642 49 969 50 587 51 041 50 771 51 858 51 516 50 242 49 818 52 010 52 852 51 270 51 813 52 267 51 058 50 967 53 229 54 652 55 052 55 620 56 421 57 005 57 399 58 714 51 999 52 676 53 169 52 902 54 327 55 300 56 711 57 979 47 376 48 123 47 984	61.539 58.491 61.581 58.535 60.078 59.763 59.542 60.698 60.701 58.721 57.632 59.181 59.037 58.296 59.284 57.729 61.593 62.766 63.681 64.337 63.600 63.274 64.132 61.838 63.819 62.668 62.271 61.445 62.473 64.651 63.592 61.491 65.033 60.965 64.487 62.947 63.730 64.446 63.736 64.413 64.423 65.699 65.297	26.423 28.940 27.846 30.360 29.799 29.464 30.810 31.656 32.874 30.716 29.848 28.697 31.402 31.729 30.654 28.858 31.024 31.766 31.998 33.005 30.917 30.954 30.271 31.549 30.141 31.493 30.725 30.439 30.725 30.439 30.725 30.439 30.725 30.439 30.725 30.439 30.725 30.439 30.725 30.439 30.725 30.74 29.940 31.072 31.233 32.311 33.100 29.847 28.996 27.542	20.00 20.00 20.00 20.00 20.00 12.86 13.82 15.06 15.36 11.50 20.00 20.00 20.00 20.00 20.00 14.37 13.98 14.72 15.56 15.10 15.01 16.69 17.26 17.79 18.91 22.03 20.00
4400	LEU	CB	48.123	64.423	29.847	15.22
4401			47.836	65.699	28.996	17.91
4402	LEU	CD1	47.984	65.297	27.542	17.36
4403		CD2	48.732	66.869	29.241	17.38
4404		Н	50.299	63.237	30.217	20.00
4405		HA	49.052	65.467	31.565	20.00
4406			47.178	63.886	29.921	20.00
4407		2HB	48.719	63.775	29.209	20.00
4408	LEU.	HG	46.811	66.008	29.196	20.00
4409	LEU	1HD1	47.280	64.485	27.341	20.00
4410	LEU	2HD1	48.978	64.923	27.304	20.00

		•		30	)9	
4411	LEU	3HD1	47.734	66.112	26.869	20.00
4412	LEU		49.761	66.547	29.286	20.00
4413	LEU		48.501	67.327	30.203	20.00
4414	LEU		48.593	67.645	28.487	20.00
4415	ALA	N	47.923	62.376	32.381	13.18
4416	ALA	CA	47.070	61.732	33.394	12.65
4417	ALA	C	47.647	61.860	34.804	14.16
4418		0		62.169	35.735	14.00
4419	ALA		46.924			13.02
	ALA	СВ	46.896	60.294	33.069	
4420	ALA	Н	48.404	61.889	31.656	20.00
4421	ALA	HA	46.109	62.245	33.400	20.00
4422	ALA	1HB	47.848	59.772	33.044	20.00
4423	ALA	2HB	46.427	60.175	32.093	20.00
4424	ALA	3HB.	46.257	59.808	33.797	20.00
4425	VAL .	N	48.993	61.669	34.953	11.78
4426	VAL	CA	49.638	61.928	36.266	12.53
4427	VAL	С	49.529	63.388	36.657	13.82
4428	VAL	0	49.106	63.654	37 <sub>.</sub> 760	12.86
4429	VAL	CB	51.060	61.470	36.199	11.66
4430	VAL	CG1	51.011	59.975	35.935	13.24
4431	VAL	CG2	51.884	61.820	37.450	9.92
4432	VAL	H	49.505	61.389	34.142	20.00
4433	VAL	HA	49.097	61.346	37.011	20.00
4434	VAL	НВ	51.525	61.950	35.339	20.00
4435	VAL	1HG1		59.680	35.046	20.00
4436	VAL		50.567	59.443	36.774	20.00
4437	VAL		52.028	59.609	35.790	20.00
4438	VAL		51.434	61.452	38.374	20.00
4439	VAL		52.023	62.899	37.543	20.00
4440	VAL		52.878	61.383	37.357	20.00
4441	ILE	N	49.859	64.316	35.726	12.29
4442	ILE	CA	49.777	65.753 ·	36.099	13.94
4443	ILE	C	48.339	66.176	36.660	14.15
4444	ILE	Ö	48.117	66.660	37.773	14.06
4445	ILE	СВ	50.394	66.654	34.959	13.50
4446	ILE	CG1	51.944	66.544	34.796	14.91
4447	ILE	CG2	50.076	68.142	35.160	12.74
4448	ILE	CD1	52.435	66.779	33.323	13.99
						20.00
4449	ILE	H	50.221	64.029	34.834	
4450	ILE .	HA	50.451	65.871	36.951	20.00
4451	ILE	HB	49.927	66.358	34.020	20.00
4452	ILE		52.222	65.531	35.081	20.00
4453	ILE		52.448	67.206	35.500	20.00
4454	ILE		50.457	68.501	36.112	20.00
	ILE		49.002	68.322	35.154	20.00
	ILE		50.501	68.767	34.372	20.00
4457	ILE		52.530	67.849	33.137	20.00
	ILE		.51.7.7.4	66.409	32.552	20.00
4459	ILE	3HD1	53.420	66.336	33.174	20.00
4460	GLU	Ν	47.359	65.839	35.881	13.27
4461	GLU	CA	45.989	66.067	36.314	12.79
4462	GLU	C,	45.607	65.346	37.640	14.22

4463 4464 4465 4466 4467 4468 4469 4470	GLU GLU GLU GLU GLU GLU GLU	O CB CG CD OE1 OE2 H HA	45.016 45.069 43.617 43.489 44.197 42.684 47.536 45.923	65.931 65.673 66.022 67.515 68.391 67.786 65.473 67.146	38.541 35.120 35.440 35.873 35.267 36.795 34.961 36.510	14.98 11.88 12.69 18.61 19.68 20.07 20.00 20.00
4471 4472 4473	GLU GLU GLU	1HB 2HB 1HG	45.179 45.379 43.215	64.633 66.212 65.380	34.870 34.239 36.227	20.00 20.00 20.00
4474 4475	GLU GLY	2HG N	42.978 46.045	65.886 64.072	34.565 37.760	20.00 14.10
4476	GLY	CA	45.654	63.221	38.876	11.35
4477 4478	GLY GLY	C	46.321 45.815	63.593 63.292	40.191 41.292	12.42 14.09
4479	GLY	H	46.512	63.723	36.945	20.00
4480	GLY.	1HA	45.949	62.203	38.620	20.00
4481 4482	GLY ALA	2HA N	44.571 47.519	63.247 64.244	38.973 40.018	20.00 12.11
4483	ALA	CA	48.374	64.823	41.082	13.24
4484	ALA	C	47.582	65.702	41.977	14.50
4485	ALA	O.	47.596	65.589	43.194 ·	15.74
4486	ALA	СВ	49.490	65.703	40.494	12.85
4487 4488	ALA ALA	H HA	47.815 48.781	64.323 63.986	39.062 41.661	20.00
4489	ALA	1HB	49.117	66.579	39.973	20.00
4490	ALA	2HB	50.058	65.139	39.757	20.00
4491	ALA	3HB	50.202	66.027	41.246	20.00
4492	LYS	Ν	46.816	66.546	41.296	15.43
4493	LYS	CA	45.724	67.345	41.899	17.97
4494	LYS	C	44.931	66.779	43.125	18.17
4495 4496	LYS LYS	O CB	44.899 44.741	67.351 67.668	44.214 40.789	16.72 17.36
4490	LYS	CG	45.473	68.507	39.735	20.89
4498	LYS	CD	44.411	69.357	39.059	19.35
4499	LYS	CE	44.941	70.273	37.899	21.45
4500	LYS	NZ	43.854	70.561	36.916	28.58
4501	LYS	Н	46.970	66.537	40.302	20.00
4502	LYS	HA	46.194	68.257	42.271	20.00
4503 4504	LYS LYS	1HB 2HB	43.929 44.281	68.238 66.774	41.230 40.403	20.00
	LYS	1HG	46.030	67.909	39.021	20.00
4506	LYS	2HG	46.191	69.182	40.190	20.00
4507	LYS	1HD	43.864	69.945	39.793	20.00
4508	LYS	2HD	43.678	68.654	38.678	20.00
4509	LYS		45.750	69.766	37.367	20.00
4510 4511	LYS. LYS	2HE 1HZ	45.330 42.957	71.199 70.859	38.323 37.339	20.00
4512	LYS	2HZ	43.637	69.601	36.519	20.00
4513	LYS	3HZ	44.134	71.121	36.095	20.00
4514	PHE	Ν	44.281	65.638	42.830	17.06

4515 4517 4518 4519 4521 4523 4523 4524 4525 4526 4527 4528 4529 4533 4533 4533 4533 4534 4534 4545 4546 4557 4558 4556 4556 4561 4562 4563 4563 4563 4563 4563 4563 4563 4563	PPPPPPPPPPPPPPPPPPILLILLILLILLILLILLILLI	2HG2 3HG2 1HD1 2HD1 3HD1 N CA C O CB CG SD CE H HA	43.646 44.714 44.482 42.997 42.385 43.226 41.030 42.767 41.444 42.892 43.750 42.235 44.244 40.363 47.496 46.963 47.403 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.734 46.963 47.634 48.913 50.089 51.391 51.905 47.175 48.874 49.285	64.836 64.518 64.738 63.581 62.573 61.617 62.660 60.863 61.903 61.951 63.045 63.045 63.045 63.045 63.045 63.045 64.058 63.619 64.058 63.619 64.058 63.619 64.058 63.820 61.500 62.830 62.830 63.820 63	43.843 44.920 46.105 43.127 44.092 44.720 44.476 45.587 46.279 41.348 42.548 42.548 43.956 43.956 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 43.715 44.366 44.361 45.366 45.367 46.363 46	17.26 17.65 17.35 16.04 18.17 17.46 18.56 17.26 16.76 20.00
4563 4564 4565	MET MET MET	1HB 2HB 1HG	49.285 48.257 50.607	68.791 68.389 67.834	46.230 44.906 44.333	20.00 20.00 20.00
4566	MET	2HG	49.687	66.363	44.396	20.00

4567	MET	1HE	51.085		68.420		47.548		20.00
4568	MET	2HE	52.319		68.603		46.291		20.00
4569	MET	3HE	52.687		67.673	•	47.749		20.00
4570	GLY	N	45.872		67.378		47.277		17.81
4571	GLY	CA	44.850		67.320		48.361		19.24
4572	GLY	C	43.474		67.982		48.080		18.12
4573 4574	GLY GLY	O H	42.634 45.672	4)	68.149 66.962		48.984 46.387		18.31
4575	GLY	п 1HA	45.072		67.841		49.231		20.00 20.00
4576	GLY	2HA	44.716		66.282		48.674		20.00
4577	ASP	N	43.290		68.369		46.814		16.53
4578	ASP	CA	41.961		68.826		46.372		17.00
4579	ASP	C	41.035		67.607	,	46.048		18.13
4580	ASP	0	40.855		67.207		44.903		19.06
4581	ASP	СВ	42.133		69.714		45.111		18.82
4582	ASP	CG	40.838		69.926		44.250		20.22
4583	ASP	OD1	39.720		69.692		44.750		19.51
4584	ASP	OD2	40.968		70.317	•	43.089		23.56
4585 4586	ASP ASP	Н .	44.012		68.210		46.144		20.00
4587	ASP	HA 1HB	41.478 42.844		69.396 69.226		47.157 44.444		20.00
4588	ASP	2HB	42.546		70.691		45.349		20.00
4589	SER	N	40.427		66.964		47.031		18.73
4590	SER	CA	39.684		65.767		46.555		16.96
4591	SER	С	38.438	1	66.178	•	45.684		18.39
4592	SER	0	37.753		65.300		45.151		18.33
4593	SER	CB	39.548		64.762		47.784		17.13·
4594	SER	OG	40.769		64.296		48.566	•	14.25
	SER		40.606	+	67.273		47.975	1116	20.00
4596	SER	HA	40.219		65.218		45.768		20.00
4597 4598	SER	1HB	38.927	:	63.897		47.443		20.00
4599	SER SER	2HB HG	38.817 41.719	1 8	65.230 64.490		48.491 48.317		20.00
4600	SER	N	38.161		67.528		45.516		17.50
4601	SER	CA ·	37.016		68.024		44.673		18.74
4602	SER	·C	37.183		67.715		43.164		17.75
4603	SER	Ο,	36.183		67.406		42.477		16.68
4604	SER	CB	36.747		69.546		44.774		19.67
4605	SER	OG	37.500		70.494		43.897		21.88
4606	SER	H	38.797		68.224		45.866		20.00
4607	SER	HA	36.131		67.483		45.019		20.00
4608	SER	1HB	36.605		69.852		45.852		20.00
4609 4610	SER SER	2HB HG	35.674 38.484	•	69.626 70.378		44.497 43.663		20.00 20.00
4611	VAL	N	38.473	*	67.708	į.	42.702		17.27
4612	VAL	CA	38.680		67.191		41.319	•	18.22
4613	VAL	C	37.910	- ,	65.970		40.883		18.73
4614		_0	37.472		65.906		39.737		19.63
4615	VAL	СВ	40.132		66.911		40.897		20.69
4616	VAL	CG1	40.884	;	66.073		41.910		17.79
4617	VAL	CG2	40.868		68.222		40.572		23.70
4618	VAL	Н	39.207		68.134		43.257		20.00

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4619	VAL	НА	38.305		67.980		40.690		20.00
4620	VAL	нв	40.169		66.379		39.947		20.00
4621	VAL		40.410		65.103		42.049		20.00
4622	VAL		40.897		66.580		42.855		20.00
4623	VAL		41.920		65.935	•	41.604		20.00
						,			
4624	VAL		41.042		68.758		41.495	.•	20.00
4625	VAL		40.304		68.883		39.916		20.00
4626	VAL		41.836		68.034		40.111		20.00
4627		N	37.827		64.997		41.788		18.28
4628	GLN	CA	37.407		63.664		41.383		19.78
4629	GLN	С	35.957		63.564		40.980		23.25
4630	GLN	Ο.	35.635		62.870		40.010		25.12
4631	GLN	CB	37.684	•	62.612		42.412		20.56
4632	GLN	CG	37.171		61.233		41.947		24.84
4633	GLN	,CD	37.628		60.156		42.928		27.26
4634	GLN	OE1	38.497		60.357		43.777	•	31.04
4635	GLN	NE2	37.005		59.010		42.735		29.42
4636	GLN	Ή	38.155		65.203		42.714		20.00
4637.	GLN	HA .	37.983		63.430		40.496		20.00
4638	GLN	·1HB	37.199		62.866		43.353	-	20.00
4639	GLN	2HB	38.752		62.567		42.609		20.00
4640	GLN	1HG	37.541		60.972		40.956		20.00
4641	GLN	2HG	36.077		61.187		41.899		20:00
4642	GLN	1HE2			58.293		43.411		20.00
4643	GLN	2HE2			58.854		42.017		20.00
4644	ASP	N ·	35.132	1	64.360		41.665		25.31
4645	ASP	CA	33.794	•	64.542		41.078		27.02
4646	ASP	C	33.714		65.515		39.851		25.79
4647	ASP	0	33.010		65.234		38.884		25.32
4648	ASP		32.728	F	64.363		42.176	**	35.78
4649	ASP	CG	32.720		62.848	:	42.170		44.92
4650	ASP	OD1	33.060		61.924		42.232		50.04
4651	ASP	OD)	31.014	ı	62.626		42.502		50.59
4652	ASP	H	35.428		64.744		42.538		20.00
4653	ASP						40.453	,	
4654	ASP	HA 1HB	33.606 31.871	• .	63.669 64.981				20.00
		1HB 2HB					41.929		20.00
4655	ASP		33.104	10	64.706		43.143		20.00
4656	GLN	N	34.626		66.532		39.807	,	23.38
4657	GLN	CA	34.903		67.171		38.484		24.32
	GLN	C	35.209		66.229		37.269		23.16
4659	GLN		34.689		66.444		36.165		21.27
4660	GLN	СВ	35.989		68.248		38.513		28.00
4661	GLN	CG	35.765		69.233		39.647	•	35.72
4662	GLN	CD	37.011		70.084		39.806		43.13
4663	GLN	OE1	37.614		70.466		38.814		48.57
4664	GLN	NE2	37.399		70.384	•	41.059		43.82
4665	GLN	H	35.105		66.754		40.660	,	20.00
4666_	_GLN	HA	.33.969.		67.668	·	_38.222_		20.00
4667	GLN	1HB	36.001		68.784		37.562		20.00
4668		2HB	36.973		67.792		38.594		20.00
4669	GLN	1HG	35.574		68.757		40.601		20.00
4670	GLN	2HG	34.922		69.879		39.419		20.00

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4671 4672 4673 4674 4675	GLN GLN TRP TRP TRP		38.245 36.969 36.067 36.319 35.089		70.918 70.145 65.207 64.264 63.487		41.061 41.925 37.463 36.366 35.918	*	20.00 20.00 20.29 19.31 19.07	
4676 4677 4678 4679 4680	TRP TRP TRP TRP TRP	O CB CG CD1 CD2	34.861 37.263 38.589 39.188 39.483		63.193 63.169 63.776 64.909 63.209		34.736 36.885 37.213 36.656 38.143		19.20 19.26 15.65 14.94 15.85	
4681 4682 4683 4684	TRP TRP TRP TRP	NE1 CE2 CE3 CZ2	40.412 40.619 39.390 41.699		65.088 64.068 62.115 63.722		37.205 38.133 38.989 38.904	, , , , t	15.54 15.86 14.86 16.78	
4685 4686 4687 4688 4689	TRP TRP TRP TRP TRP	CZ3 CH2 H HA 1HB	40.479 41.631 36.508 36.722 37.452		61.761 62.585 65.153 64.799 62.427		39.786 39.740 38.360 35.513 36.116		14.37 15.56 20.00 20.00 20.00	
4690 4691 4692 4693 4694	TRP TRP TRP TRP TRP	2HB HD1 HE1 HE3 HZ2	36.849 38.745 41.059 38.490 42.571		62.663 65.543 65.811 61.508 64.373		37.759 35.900 37.001 38.989 38.864		20.00 20.00 20.00 20.00 20.00	
4695 4696 4697 4698 4699	TRP TRP LYS LYS LYS	HZ3 HH2 N CA C	40.436 42.485 34.338 33.082 32.154	. • •	60.909 62.328 63.136 62.421 63.264		40.460 40.349 36.976 36.730 35.758	: Y : - +	20.00 20.00 21.19 24.17 25.28	
4700 4701 4702 4703 4704	LYS LYS LYS LYS LYS	O CB CG CD CE	31.736 32.453 31.329 30.909 29.764		62.892 62.137 61.095 60.906 59.913		34.656 38.108 38.057 39.490 39.692		26.15 26.43 32.00 38.87 45.47	
4705 4706 4707 4708	LYS LYS LYS	NZ H HA 1HB	29.279 34.648 33.334 32.101		60.040 63.302 61.452 63.055		41.099 37.917 36.303 38.569		51.74 20.00 20.00 20.00	
4710 4711 4712 4713	LYS LYS LYS LYS LYS	2HD	33.239 31.674 30.495 30.615 31.768	÷	61.739 60.151 61.432 61.873 60.576	(¥)	38.752 37.631 37.435 39.902 40.078		20.00 20.00 20.00 20.00 20.00	
4716 4717	LYS LYS LYS LYS -LYS	1HE 2HE 1HZ 2HZ -3HZ	30.136 28.947 29.033 30.103 -28.496	· ·	58.898 60.090 61.040 59.869 59.397		39.499 38.984 41.269 41.719 41.317		20.00 20.00 20.00 20.00 20.00	
4719 4720 4721 4722	GLU	N CA C O	32.050 31.455 32.006 31.332	. *	64.521 65.533 65.577 65.367	;	36.209 35.365 33.869 32.866	2	26.22 27.87 26.53 30.42	İ

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4723	GLU	СВ	31.542		66.807		36.263		34.80
4724	GLU	CG .	30.693		66.725		37.585		47.61
4725	GLU	CD	29.211		66.293		37.462		55.88
4726	GLU	OE1	28.422		67.129		36.998		59.76
4727	GLU	OE2	28.883		65.134		37.826		61.03
4728		H	32.243		64.712	,	37.176		20.00
4729	GLU	HA	30.410		65.251		35.271		20.00
4730	GLU	1HB	31.181		67.661		35.698		20.00
4731	GLU	2HB	32.564		67.009		36.547	,	20.00
4732	GLU	1HG			67.672		38.117		20.00
4733	GLU	2HG	31.080		65.993		38.276		20.00
	LEU	N	33:317		65.813	٠	33.793	:	24.19
4735	LEU	CA ·	34.088		66.004		32.540		23.43
	LEU	C	34.108		64.773	•	31.610	<b>√</b> 11	24.08
4737		Ö	34.441		64.885		30.447		22.85
4738	LEU	CB	35.553		66.277	, i	32.956		25.02
4739	LEU	CG	35.989		67.725	;	32.878		25.39
4740	LEU	CD1	37.358		67.805	•	33.605	•	27.03
4741	LEU	CD2	34.947	:.	68.615		33.583		28.40
4742	LEU	Н	33.756	•	65.871		34.690		20.00
	LEU	НА	33.699		66.832		31.949		20.00
4744	LEU	1HB	36.274		65.707	٠.	32.372	· ·	20.00
4745	.LEU	2HB	35.693	٠.	65.929		33.976		20.00
4746	LEU	HG	36.089		68.033		31.838	1	20.00
4747	LEU	1HD1	38.106		67.202		33.097		20.00
4748	LEU	2HD1	37.284		67.450		34.634		20.00
4749	LEU	3HD1	37.742	,	68.825	. , '	33.641		20.00
4750	LEU	1HD2	34.769		68.279	•	34.605		20.00
	LEU	4 4 11 1440 1 1	33.982		68.644		33.078		20.00
4752	LEU		35.307		69.642		33.676		20.00
4753	SER	N	33.842		63.591		32.146		23.22
4754	SER	CA	34.023		62.430		31.279		24.53
4755		C	32.729		62,144	,	30.475	•	26.87
	SER	0	,		61.539	1	29.414		26.47
4757		CB	34.168		61.268	100	32.238		23.92
4758	SER	OG.	32.851		60.942		32.814		27.33
4759	SER	H	33.548		63.568		33.105		20.00
4760		HA	34.900		62.512		30.638		20.00
4761		1HB	35.074		61.383		32.907		20.00
4762	SER		34.520		60.416		31.633	· · .	20.00
4763		HG	32.219		61.568		33.301	. ', 1;	20.00
	HIS	N ·	31.582		62.546		31.085	• •	29.80 33.13
4766	HIS	CÁ C	30.257 29.778		62.150 60.651		30.557 30.861		33.64
4767		0	29.176		59.996		29.999		31.30
4768		CB	30.150		62.548		29.063	٠.	36.51
4769		CG.	30.638		63.959		28.940		40.71
4770	HIS	ND1	31.746		64.287		28.246		43.97
4771	HIS	CD2	30.108		65.137		29.519		42.68
4772		CE1	31.894		65.632		28.386		43.60
4773		NE2	30.917		66.174	• • • • • •	29.151		42.67
4774	HIS	Н	31.682		63.199		31.841		20.00
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HIS	НА	29.569	62.773	31.127	20.00
HIS	1HB	29.124	62.477	28.704	20.00
HIS	2HB	30.767	61.968	. 28.391	20.00
HIS	HD1	32.351	63.667	27.792	20.00
HIS	HD2	29.244	65.194	30.166	20.00
HIS	HE1	32.696	66.220	27.957	20.00
GLU	Ν	30.038	60.148	32.122	35.19
		29.812	58.725	32.461	36.70
					38.60
			and the second s		37.53
					35.82
					36.20
					38.96
					36.60
					41.14
					20.00
			••		20.00
					20.00
					20.00
					20.00
					42.63
					46.40
			and the second s		47.19
ASP	Ö		· ·	30.242	46.38
ASP	CB	25.156	59.121	33.508	50.98
ASP	CG	25.757	58.386	34.723	57.85
ASP	OD1	25.724	57.146	34.731	60.50
ASP.	. OD2 -	26.292	59.021	35.648	61.02
ASP	OXT	24.582	58.276	30.208	48.57
		27.519		32.926	20.00
					20.00
					20.00
					20.00
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					0.00
					0.00
					0.00
OC_	O13	50.068	42.623	14.140	0.00
OC <u>-</u>	014	52.998_	39.925	16.291	0.00
OC_	015	51.718	37.289	15.246	0.00
OC_	H16	51.994	36.801	14.488	0.00
OC_	C17	52.602	42.662	15.945	0.00
OC_	C18	51.989	43.758	16.830	0.00
	HHHHGGGGGGGGGGGGGGAAAAAAAAAAOOOOOOOOOOO	HIS 2HB HIS HD1 HIS HD2 HS HD1 HS HD1 HS HD1 HS HD1 HS HD1 HS HD1 HS HD1 HS HD2 HS HD2 HS HD1 HS HD2 HS HS HS HS HS HS HS HS HS br>HS HS HS HS HS HS HS HS HS HS HS HS HS HS HS HS HS HS H	HIS 1HB 29.124 HIS 2HB 30.767 HIS HD1 32.351 HIS HD2 29.244 HIS HE1 32.696 GLU N 30.038 GLU CA 29.812 GLU C 28.365 GLU CB 30.385 GLU CB 30.385 GLU CD 30.185 GLU CD 30.185 GLU OE1 31.324 GLU H 30.309 GLU HA 30.412 GLU 1HB 31.453 GLU 2HB 30.305 GLU 1HG 28.617 GLU 2HG 29.808 ASP N 27.355 ASP CA 25.947 ASP CA 25.947 ASP CB 25.156 ASP CB 25.156 ASP CB 25.757 ASP OD1 25.724 ASP OD2 26.292 ASP OXT 24.582 ASP H 27.519 ASP HA 25.950 ASP 1HB 24.118 ASP 2HB 25.188 OC_ C1 49.640 OC_ C2 50.787 OC_ C3 51.008 OC_ C4 50.154 OC_ C5 49.053 OC_ C6 48.782 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C10 52.036 OC_ C17 52.602	HIS 1HB 29.124 62.477 HIS 2HB 30.767 61.968 HIS HD1 32.351 63.667 HIS HD2 29.244 65.194 HIS HE1 32.696 66.220 GLU N 30.038 60.148 GLU CA 29.812 58.725 GLU C 28.365 58.293 GLU O 28.187 57.171 GLU CB 30.385 58.171 GLU CG 29.685 58.624 GLU CD 30.185 57.959 GLU OE1 31.324 57.527 GLU OE2 29.434 57.837 GLU H 30.309 60.799 GLU HA 30.412 58.167 GLU 1HB 31.453 58.377 GLU 2HB 30.305 57.095 GLU 1HG 28.617 58.448 GLU 2HG 29.808 59.700 ASP N 27.355 59.161 ASP CA 25.947 58.715 ASP C 25.333 59.095 ASP O 25.663 60.179 ASP CB 25.156 59.121 ASP CG 25.757 ASP CB 25.156 59.121 ASP CG 25.757 ASP OD2 26.292 59.021 ASP ONT 24.582 58.276 ASP H 27.519 60.002 ASP HA 25.950 57.622 ASP 1HB 24.118 58.815 ASP 2HB 25.188 60.198 OC_ C1 49.640 37.719 OC_ C2 50.787 38.141 OC_ C3 51.008 39.495 OC_ C1 49.640 37.719 OC_ C2 50.787 38.141 OC_ C3 51.008 39.495 OC_ C1 49.640 37.719 OC_ C2 50.787 38.141 OC_ C3 51.008 39.495 OC_ C1 49.640 37.719 OC_ C6 48.782 38.701 OC_ TH 49.187 36.718 OC_ C10 52.036 40.047 OC_ C6 48.782 38.701 OC_ TH 49.187 36.718 OC_ C10 52.036 40.299 OC_ N11 51.728 41.572 OC_ C12 50.628 41.640 OC_ O13 50.068 42.623 OC_ O14 52.998 39.925 OC_ O15 51.718 37.289 OC_ O15 51.718 37.289 OC_ H16 51.994 36.801 OC_ C17 52.602 42.662	HIS 1HB 29.124 62.477 28.704 HIS 2HB 30.767 61.968 28.391 HIS HD1 32.351 63.667 27.792 HIS HD2 29.244 65.194 30.166 HIS HE1 32.696 66.220 27.957 GLU N 30.038 60.148 32.122 GLU CA 29.812 58.725 32.461 GLU C 28.365 58.293 32.195 GLU O 28.187 57.171 31.764 GLU CB 30.385 58.171 33.820 GLU CG 29.685 58.624 35.121 GLU CD 30.185 57.959 36.421 GLU OE1 31.324 57.527 36.458 GLU OE2 29.434 57.837 37.399 GLU H 30.309 60.799 32.826 GLU H 30.309 60.799 32.826 GLU H 30.309 60.799 32.826 GLU HB 31.453 58.377 33.892 GLU 1HB 31.453 58.377 33.892 GLU 2HB 30.305 57.095 33.743 GLU 1HG 28.617 58.448 35.059 GLU 2HG 29.808 59.700 35.259 ASP CA 25.947 58.715 32.232 ASP CA 25.947 58.715 32.232 ASP CA 25.947 58.715 32.232 ASP CB 25.156 59.121 33.508 ASP CB 25.156 59.121 33.508 ASP CB 25.757 58.386 34.723 ASP CB 25.757 58.386 34.723 ASP CD 25.663 60.179 30.242 ASP CB 25.757 58.386 34.723 ASP OD1 25.724 57.146 34.731 ASP OD2 26.292 59.021 35.648 ASP OD2 26.292 59.021 35.648 ASP OD3 24.582 58.276 30.208 ASP HA 25.950 57.622 32.219 ASP HB 24.118 58.815 33.431 ASP 2HB 25.188 60.198 33.649 OC_ C1 49.640 37.719 14.003 OC_ C2 50.787 38.141 14.736 OC_ C3 51.008 39.495 14.916 OC_ C4 50.154 40.364 14.322 OC_ C5 49.053 40.047 13.581 OC_ C6 48.782 38.701 13.402 OC_ TH 49.840 40.820 13.123 OC_ C1 52.036 40.299 15.636 OC_ N11 51.728 41.572 15.438 OC_ C12 50.628 41.640 14.622 OC_ O14 52.998 39.925 15.469 OC_ C14 50.154 40.820 13.123 OC_ C15 51.718 37.289 15.246 OC_ C16 52.036 40.299 15.636 OC_ N11 51.728 41.572 15.438 OC_ C19 52.036 40.299 15.636 OC_ O15 51.718 37.289 15.246 OC_ O14 52.998 39.925 16.291 OC_ O15 51.718 37.289 15.246 OC_ O15 51.718 37.289 15.246 OC_ O15 51.718 37.289 15.246 OC_ O14 52.998 39.925 16.291 OC_ O15 51.718 37.289 15.246 OC_ O14 52.998 39.925 15.6291 OC_ O15 51.718 37.289 15.246

19	OC_	H19	53.162		43.136		15.129		0.00	
20	OC_	H20	53.441		42.237		16.493		0.00	
21	OC_	021	52.823	•	43.764		17.935		0.00	
22	OC_	C22	52.911		44.965		18.630		0.00	
23	00_	C23	51.464		45.106		19.104		0.00	
24 25	00_	C24	50.401		44.489		18.575		0.00	
26	OC_	C25 6H2	50.577 52.083		43.599 44.717		17.383 16.324		0.00	
27	OC_	H27	53.299		45.778		18.011		0.00	
28	OC_	H28	53.618		44.846		19.458		0.00	
29	OC_	9H2	49.817		43.779	•	16.639		0.00	
30	OC_	0H3	50.512		42.573		17.745		0.00	
31	OC_	S31	50.995		45.860		20.622		0.00	
32	OC_	C32	49.309		45.615		20.276		0.00	
33	OC_	C33	49.082		44.948		19.126		0.00	
34	OC_	N34	48.236		46.161		21.034		0.00	
35	OC_	C35	48.405		46.833		22.220		0.00	
36 37	OC_	C36	47.021		47.267		22.889		0.00	
3 <i>1</i>	OC_	O37 O38	46.749 46.116		48.354 46.291		23.338		0.00	
39	OC_	O39	49.477		47.023		23.042 22.820		0.00	
40	OC_	C40	47.697		44.874		18.500		0.00	
41	OC_	041	46.650		44.969		19.070		0.00	
42	OC	042	47.614		44.562		17.174		0.00	
43		3H4	46.996		43.859		17.092		0.00	
44	oc_	H44	45.261		46.650		23.184		0.00	
45	OC_	H45	47.336		46.168		20.596		0.00	
1	TIP	OH2	55.419		44.829		16.389		20.00	
2 .	TIP	H1 .	55.536		44.889		17.342		20.00	
3	TIP	2H	55.300		45.765		16.164		20.00	
4 5	TIP	OH2 1H	50.936 51.119		38.099 38.023		22.176		20.00	
6	TIP	H2	50.913	•	39.042		23.113 22.012		20.00 20.00	
7	TIP	OH2	29.774		30.704		38.242		20.00	
8	TIP	1H	29.956		30.628		39.179		20.00	
9	TIP	H2	29.750		31.647		38.078		20.00	
10	TIP	OH2	45.277		35.890		28.823		20.00	
11	TIP	1H	45.460		35.813		29.759		20.00	
12	TIP	H2	45.253		36.832		28.659		20.00	
13	TIP	OH2	58.027		40.785		28.285		20.00	
14	TIP	1H	58.210		40.709		29.222		20.00	
15 16	TIP	H2	58.004		41.728		28.121		20.00	
16 17	TIP TIP	OH2 H1	40.267 40.450		36.083		19.326		20.00	
18	TIP	2H	40.450		36.007 37.026		20.263 19.162		20.00	
19	TIP	OH2	53.647		32.258		38.649		20.00	
20	TIP	1H	53.830		32.182		39.585		20.00	
21	TIP	H2	53.623		33.200		38.484		20.00	
22	TIP	OH2	48.317	:	32.381		26.654	• • • •	20.00	-
23	TIP	1H	48.499		32.305		27.591		20.00	
24	TIP	H2	48.293		33.324		26.490		20.00	
25	TIP	OH2	38.532		50.364		24.358		20.00	

				*					
26	TIP	1H	38.714	50	0.288	2	5.294		20.00
27	TIP	H2	38.508		.307		4.194		20.00
28	TIP	OH2	43.205		2.135		2.424		20.00
29	TIP	1H	43.387	_	2.059		3.360		20.00
30	TIP	2H	43.181	43	3.078	4	2.260		20.00
31	TIP	OH2	38.345	49	9.997	2	1.607		20.00
32	TIP	H1	38.528	49	9.921	2	2.543		20.00
33	TIP	H2	38.321		.940		1.443		20.00
34	TIP	OH2	48.352		0.997		7.771		20.00
35	TIP	1H	48.535		).921		8.708		20.00
36	TIP	H2	48.329		.940		7.607	,	20.00
37	TIP	OH2	48.526		1.351		3.768		20.00
38	TIP	H1	48.709		1.275		4.705		20.00
39	TIP	2H	48.502	- 25	5.294	2	3.604		20.00
40	TIP	OH2	30:895	32	2.557	4	9.007		20.00
41	TIP	H1	31.078	32	2.480	4	9.944		20.00
42	TIP	2H	30.871	33	3.499	4	8.843	•	20.00
43	TIP	OH2	48.519		0.061		1.813	*	20.00
44	TIP	H1	48.702		9.985		2.750		20.00
45	TIP	2H	48.495		.003		1.649		20.00
46	TIP	ÖH2	57.848		.344		2.042		20.00
47	TIP	H1	58.031		.268		2.978	,	20.00
48	TIP	2H	57.825		2.287		1.877		20.00
49	TIP	OH2	54.834		5.583		1.192		20.00
50 ·	TIP	1H	55.017		5.507		2.129	٠,	20.00
51	TIP	H2	54.811	36	5.525	2	1.028		20.00
52	TIP	OH2	21.604	40	.670	3	7.071		20.00
53	TIP	H1	21.787	40	.594	3	8.007		20.00
54	TIP .	2H	21.581	41	.613		6.907		20.00
55	TIP	OH2	61.252		.808		7.483		20.00
56	TIP	H1	61.435		.732		8.420		20.00
57	TIP	2H	61.229		3.751		7.319		20.00
58	TIP	OH2	66.912		.122		0.151		20.00
59		H1	67.094		0.046		1.088		20.00
	TIP.								
60 -	TIP	2H	66.888		0.065		9.987		20.00
61	TIP	OH2	23.155		.413		5.818		20.00
62	TIP	H1	23.337		3.337		6.755		20.00
63	TIP	2H	23.131		355		5.654		20.00
64	.TIP	OH2	52.477	58	3.511		8.314		20.00
65	TIP	H1 ,	52.659	58	.434	19	9.250		20.00
66	TIP	2H	52.453	59	.453	18	8.150		20.00
67	TIP	OH2	33.877	44	.186	2	3.766		20.00
68	TIP	1H	34.060		.110		4.702		20.00
69	TIP	H2	33.853		.129		3.602		20.00
70	TIP	OH2	36.071		3.377		8.280		20.00
71	TIP	H1	36.254		3.301		9.216		20.00
72	TIP	2H			.320		8.116		20.00
			36.047						
-73	TIP_	OH2	57.951		2.393_		2.291	· · -	20.00
74	TIP	1H	58.133		2.317		3.228		20.00
75	TIP	H2	57.927		.335	-1	2.127	,	20.00
76	TIP	OH2	43.946		.374		4.700		20.00
77	TIP	1H	44.128	30	.298	4	5.637		20.00

	•					5.15	9		
78	TIP	H2	43.922		31.316		44.536	_	20.00
79	TIP	OH2	23.284		48.767		33.067	-	20.00
80	TIP	1H	23.466		48.691		34.004		20.00
81	TIP	H2	23.260		49.710		32.903		20.00
82	TIP	OH2	34.465	• -	36.411		46.836		20.00
83	TIP	1H	34.648		36.335		47.773		20.00
84									20.00
	TIP	H2	34.441		37.353		46.672		
85	TIP	OH2	47.183		59.524		19.471		20.00
86	TIP	H1	47.365		59.448		20.407		20.00
87	TIP	2H	47.159		60.467		19.307		20.00
88	TIP	OH2	38.194		26.639		27.880		20.00
89	TIP	1H .	38.377		26.563		28.816		20.00
90	TIP	H2	38.170		27.581		27.716		20.00
91	TIP	OH2	63.749		46.405		40.207		20.00
92	TIP	1H	63.932		46.329		41.143		20.00
93	TIP	H2	63.726	,	47.347		40.043		20.00
94	TIP	OH2	38.952	,	29.220	•	51.044		20.00
95	TIP	1H	39.135		29.144		51.980		20.00
96	TIP	H2	38.928		30.162		50.880		20.00
97	TIP	OH2	22.585		40.880		29.562		20.00
98	TIP	H1	22.768		40.804		30.498		20.00
99	TIP	2H	22.562		41.823		29.398		20.00
100	TIP	OH2	60.690		27.339		33.408		20.00
101	TIP	1H	60.873		27.263		34.345		20.00
102	TIP	H2	60.666		28.282		33.244	•	20.00
103	TIP	OH2	44.387		24.820		39.848	•	20.00
104	TIP	H1	44.570		24.744		40.784		20.00
105	TIP	H2	44.363		25.763		39.684		20.00
106	TIP	OH2	47.685		57.349		44.874		20.00
107	TIP	1H	47.867	•	57.272		45.810		20.00
108	TIP	H2	47.661	* .	58.291		44.710		20.00
109	TIP	OH2	67.071		45.345		34.784		20.00
110	TIP	H1	67.254		45.268		35.720		20.00
111	TIP	2H	67.047		46.287		34:620		20.00
112	TIP	OH2	45.116		59.190		18.168		20.00
113	TIP	H1	45.298		59.114	•	19.105		20.00
114	TIP	2H	45.092		60.133		18.004		20.00
115	TIP	OH2	60.283		64.299		18.011		20.00
116	TIP	H1	60.466		64.223	•	18.947		20.00
	TIP								20.00
117		2H	60.259	٠.	65.241		17.847		
118	TIP	OH2	60.415		30.584		33.261		20.00
119	TIP	H1 -	60.598		30.508	ė	34.198		20.00
120	TIP	2H.	60.392		31.527		33.097		20.00
121	TIP	OH2	60.024	•	47.287		40.698		20.00
122	TIP	1H ·	60.207		47.211		41.634	• .	20.00
123	TIP	H2	60.001		48.230		40.534		20.00
124	TIP	OH2	37.196	,	`38.650		46.459		20.00
125	TIP	H1	37.379		38.574		47.395		20.00
126	TIP	2H	37.172		39.592		46.294		20.00
127	TIP	OH2	46.215		64.400		22.087		20.00
128	TIP	H1	46.398		64.324		23.024		20.00
129	TIP	2H	46.191		65.343		21.923		20.00

130 131 132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147 148 149 150 151 153 154 155 156 157 158 160 161 162 163 164 165 166 166 166 166 166 166 166 166 166		OH2 H1 2H 2H 2H 2H 2H 2H 2H 2H 2H 2H 2H 2H 2H	32.296 32.479 32.272 25.133 25.316 25.109 63.940 64.123 63.917 42.953 43.135 42.929 31.728 31.910 31.704 63.074 63.074 63.074 63.256 63.050 57.929 58.112 57.906 37.261 37.444 37.238 49.491 49.673 49.467 58.235 58.417 58.211 39.581 39.581 39.587 26.827 26.827	42.095 42.019 43.038 25.020 24.944 25.962 65.238 65.162 66.181 24.320 24.244 25.263 20.196 20.120 21.139 44.498 44.422 45.441 49.570 49.494 50.513 57.254 58.273 44.986 44.910 45.929 25.415 25.339 26.358 24.265 24.189 25.208 29.789	24.175 25.112 24.011 39.586 40.523 39.422 29.552 30.489 29.388 36.755 37.692 36.591 39.928 40.864 43.600 42.500 22.490 23.426 22.325 21.133 22.070 20.969 44.949 45.886 44.785 34.562 35.498 34.398 39.192 40.129 39.028 39.722 40.658 25.558	20.00 20.00
163	TIP	OH2	26.644 26.827 26.620 46.323 46.506	29.865	39.722	20.00 20.00 20.00 20.00 20.00
100	1 11	411	46.300	77.701	17.701	20.00



Table of the orthogonal three dimensional coordinates in Ångstroms and B factors ( $Å^2$ ) for Protein Tyrosine Phosphatase 1B complexed with 2-(oxalyl-amino)-7-(1,1,3-trioxo-1*H*-benzo[d]isothiazol-3-yloxomethyl)-4,7-dihydro-5*H*-thieno[2,3-c]pyran-3-carboxylic acid and the water molecule which forms hydrogen bonds with the pyran oxygen atom, the side chain oxygen atom and aspartic acid 48 (Example XX).

No	Amino	acid	X	Υ	Z	В
1	GLU.	Ν.	21.703	70.016	37:889	44.68
2	GLU	CA	20.473	69.206	37.782	
3	GLU	C .	20.313		36.438	42.17
4	GLU	0	20.963	68.696		41.34
5	GLU -	CB	19.333	70.227	37.986	46.14
6	GLU	CG	17.913	69.694	38.169	53.54
7	GLU	CD	17.723	68.479	39.088	61.73
8	GLU .	OE1	16.693	68.363	39.735	64.31
9	GLU	OE2.	18.581	67.618	39.170	63.66
10	MET	N	19.366	67.432	36.416	39.32
11	MET	CA	18.893	66.684	35.226	34.55
12	MET	С	18.088		34.297	32.30
13	MET	0	18.046	67.444		32.26
14	MET	CB	17.971	65.501	35.627	34.25
15	MET	CG		65.884		32.80
16 .	MET	SD	15.642		36.918	29.12
17	MET GLU	CE	16.780	63.485	37.833	28.64 32.02
18	GLU	N- CA	17.444 16.660	68.545 69.568	.34.943 34.282	35.48
19	GLU	C	17.565	70.545	33.478	34.80
20 21	GLU	0	17.303	70.861	32.328	34.07
22	GLU	СВ	15.843	70.226	35.377	37.66
23	GLU	CG	14.638	71.016	34.859	41.16
24	GLU	CD	13.640	71.228		
25	GLU	OE1	14.018	71.066		43.98
26	GLÜ	OE2	12.488	71.540	35.734	
<b>27</b> '	LYS	N	18.704	70.909	34.098	35.30
28	LYS	CA	19.749	71.591	33.318	36.11
29	LYS	C	20.297	70.750	32.115	33.14
30	LYS	Ō	20.337	71.206	30.978	32.17
31	LYS	СВ	20.887	72.023	34.258	40.81
32	LYS	CG	20.524	73.209	35.174	47.76
33	LYS	CD	21.621	73.495	36.226	53.62
-34	LYS	· CE	21.051	73.908	-37-587	-55:86-
35	LYS	NZ	21.885	73.368	38.677	56.39
36	GLU	N -		69.500		
37	GLU	CA		68.597		
38	GLU	С		68.489		
39	GLU	0	20.353	68.592	29.020	29.85

40 41 42 43 44 45 46 47 48 49 50 51 52 53 55 55 57 58 59	GLU GLU GLU GLU GLU PHE PHE PHE PHE PHE PHE PHE GLU GLU	CB CCD OE1 OE2 N CA C O CB CCD1 CCE2 CZ N CA CO	21.406 67.217 31.894 32.53 21.384 66.134 30.814 37.49 22.178 64.888 31.173 38.32 22.206 64.433 32.300 42.27 22.785 64.379 30.267 40.62 18.794 68.321 30.587 30.73 17.727 68.276 29.603 32.45 17.750 69.480 28.642 34.85 17.904 69.288 27.453 33.64 16.381 68.223 30.320 30.35 15.294 67.870 29.350 26.13 15.021 66.538 29.085 25.00 14.582 68.860 28.688 22.62 14.068 66.173 28.147 24.45 13.636 68.508 27.740 23.14 13.391 67.166 27.459 23.15 17.650 70.698 29.207 38.88 17.603 71.927 28.405 42.49 18.812 71.977 27.485 40.63 18.782 72.264 26.313 38.60
61 62 63 64 65	GLU GLU GLU GLN	CG CD OE1 OE2 N	16.586 73.249 30.307 59.10 15.360 73:806 29.606 67.27 15.504 74.386 28.522 72.82 14.274 73.680 30.153 69.58 19.926 71.681 28.084 38.72
66 67 68 69	GLN GLN GLN	CA C O CB	21.182 71.677 27.384 39.05 21.295 70.629 26.263 36.85 21.920 70.876 25.239 37.31 22.160 71.459 28.488 43.39
70	GLN	CG	23.600 71.228 28.046 51.02
71	GLN	CD	24.424 70.842 29.287 57.42
72	GLN	OE1	25.562 70.436 29.224 62.88
73	GLN	NE2	23.787 70.955 30.441 59.65
74	ILE	N	20.644 69.479 26.488 33.52
75	ILE	CA	20.568 68.460 25.480 28.82
76	ILE	C	19.654 68.921 24.295 27.31
77	ILE	O	19.969 68.894 23.110 27.61
78	ILE	CB	20.239 67.069 25.944 27.67
79	ILE	CG1	21.327 66.546 26.867 25.62
80	ILE	CG2	20.110 66.093 24.775 24.46
81	ILE	CD1	20.897 65.251 27.523 28.44
82	ASP	N	18.496 69.346 24.737 27.29
83	ASP	CA	17.539 69.884 23.816 28.22
84 85 86 87	ASP ASP ASP ASP	C O CB CG	17.959 09.004 23.016 20.22 18.093 71.055 22.974 29.77 17.950 71.106 21.763 29.46 16.329 70.307 24.644 26.78 15.094 69.454 24.346 25.32
88	ASP	OD1	15.182 68.402 23.784 25.41 14.023 69.884 24.659 25.96 18.772 71.968 23.654 34.04 19.284 73.163 23.027 38.17
89	ASP	OD2	
90	LYS	N	
91	LYS	CA	

92 93 94 95 96 97 98 99 100 101 103 104 105 107 108 109 110 111 113 114 115 116 117 118 119 120 121 123 124 125 126 131 131 131 131 131 131 131 131 131 13	LYSSSSSSSSSSSSSSSSSTTTTTTTTTTTTTAAAAAAAAA	COCGCCNNCCOONCCOONCCOORCCOCCCNCCOONCCOORC	20.201 72.782 21.859 38.28 20.020 73.189 20.713 41.49 20.015 74.041 24.063 43.91 19.071 74.819 25.012 52.90 19.799 75.829 25.904 58.71 18.834 76.499 26.900 60.27 19.587 77.453 27.717 61.96 21.163 71.912 22.177 35.28 22.098 71.545 21.090 33.50 21.511 70.377 20.261 33.89 22.246 69.714 19.554 35.89 23.342 70.937 21.788 32.34 22.978 69.903 22.780 34.77 20.201 70.097 20.417 32.04 19.593 68.985 19.695 31.04 20.362 67.670 19.728 31.12 20.378 66.986 18.724 34.31 20.977 67.282 20.858 29.06 21.777 66.031 20.725 26.56 21.199 64.762 21.398 25.00 21.940 63.835 21.667 26.10 23.233 66.335 21.115 28.79 23.361 67.063 22.382 32.58 19.855 64.639 21.554 22.56 19.293 63.388 22.093 19.31 19.786 62.100 21.381 18.55 20.083 61.102 22.013 20.55 17.751 63.440 22.080 17.74 17.247 64.345 23.180 17.53 16.779 65.654 23.022 14.15 17.312 64.085 24.605 16.03 16.593 66.177 24.251 16.66 16.887 65.268 25.251 17.66 17.675 63.004 25.350 11.39 16.898 65.347 26.623 18.08 17.676 63.059 26.745 10.09 17.279 64.235 27.383 15.36 19.871 62.143 20.056 17.72 20.202 60.910 19.307 17.15 21.652 60.428 19.547 15.94 21.895 59.243 19.607 16.09 20.042 61.212 17.815 15.04 22.583 61.369 19.645 15.86 23.953 61.087 20.057 16.56 24.632 59.609 21.781 18.09 24.761 62.357 19.906 14.93
136	ALA	C	24.062 60.639 21.520 17.32
137	ALA	0	24.632 59.609 21.781 18.09
139	ILE	N	23.417 61.377 22.433 17.87
140		CA	23.245 60.832 23.789 19.03
141	ILE	C	22.763 59.345 23.796 18.11
142	ILE	O	23.305 58.438 24.409 18.57
143	ILE	CB	22.216 61.696 24.560 21.02

146 147 149 151 153 154 156 157 158 161 163 164 165 167 169 171 173 174 175 177 178 179 179 179 179 179 179 179 179 179 179	TTTTTTTTTTTGGGGGGGGGAAAAAAAIIILLLLLLEGGGGGGGGGGTTTTTTTTTTTTGGGGGGGGGG	N C C O C C C C C O N C C O C C C O N N C C O C C O D N C C O C C C D N C C O C C C C N C C O C C C C C N C C O C C C C	21.120 57.794 23.060 17.24 22.043 56.788 22.376 19.36 22.300 55.706 22.907 19.20 19.754 57.851 22.383 15.19 19.119 56.501 22.280 15.92 18.853 55.782 23.434 14.65 18.790 55.967 21.035 16.31 18.267 54.556 23.417 13.96 18.159 54.733 20.979 14.00 17.905 54.008 22.163 14.32 17.345 52.759 21.944 11.00 22.561 57.160 21.194 21.20 23.605 56.331 20.588 24.13 24.713 56.001 21.593 21.45 25.184 54.880 21.631 20.37 24.248 56.982 19.353 34.41 25.321 56.035 18.773 49.84 26.136 56.556 17.565 62.54 26.977 55.883 16.989 66.64 25.813 57.794 17.172 67.57 25.085 57.002 22.391 21.19 26.174 56.832 23.339 23.68 25.858 55.716 24.328 23.38 26.600 54.749 24.489 24.87 26.451 58.140 24.079 28.19 27.103 59.205 23.199 31.40 27.483 58.905 22.061 34.52 27.221 60.349 23.646 34.31 24.627 55.828 24.873 21.79 24.154 54.745 25.716 20.54 24.220 53.381 24.999 20.53 24.664 52.384 25.562 19.96 22.722 55.057 26.188 19.90 22.746 56.121 27.295 18.92 22.002 53.809 26.709 13.13 21.427 56.904 27.348 23.15 23.711 53.345 23.741 20.41 23.715 52.099 22.985 22.09 25.141 51.469 22.930 23.43 25.286 50.258 23.023 21.33 23.088 52.300 21.584 24.59 21.532 52.302 21.526 27.48
190	ARG	-CD	20.912-52.327-20.101-32.84-19.450 52.168 20.157 43.35 18.645 51.724 19.167 44.57 19.112 51.575 17.953 46.24
191	ARG	NE	
192	ARG	CZ	
193	ARG	NH1	
194	ARG	NH2	17.395 51.443 19.393 38.15
195	HIS	N	26.156 52.353 22.819 26.24

196	HIS	CA	27.538 51.876 22.706 29.54
197	HIS	C	28.137 51.358 24.001 30.14
198	HIS	0	28.822 50.348 24.027 31.41
199	HIS	CB	28.432 52.975 22.150 35.11
200 201	HIS HIS	CG ND1	28.242 52.899 20.676 45.77 28.436 51.769 19.968 50.82
202	HIS	CD2	27.726 53.883 19.830 48.72
203	HIS	CE1	28.026 52.047 18.726 53.20
204	HIS	NE2	27.594 53.318 18.613 51.96
205	GLU	N	27.850 52.116 25.059 29.74
206 207	GLU GLU	CA C	28.217 51.750 26.423 26.86 27.461 50.503 26.956 25.10
208	GLU	Ö	27.958 49.784 27.824 25.66
209	GLU	СВ	27.885 52.987 27.258 29.73
210	GLU	CG	28.749 54.190 26.816 36.62
211 212	GLU	CD OF1	28.227 55.576 27.238 42.17
212	GLU GLU	OE1 OE2	27.415 55.678 28.155 43.56 28.663 56.549 26.634 42.82
214	ALA	N	26.224 50.276 26.460 22.66
215	ALA	CA	25.424 49.170 27.037 19.17
216	ALA	C	26.192 47.808 27.064 17.78
217	ALA	O	27.027 47.515 26.229 18.91
218 219	ALA SER	CB N	24.115 49.032 26.262 11.88 25.866 47.002 28.076 16.59
220	SER	CA	26.466 45.670 28.261 15.02
221	SER	C	26.067 44.661 27.239 15.21
222	SER	0	25.057 44.724 26.572 14.30
223	SER	CB OG	25.928 45.219 29.658 13.45 26.076 46.179 30.730 20.52
224 225	SER	N	26.076 46.179 30.730 20.52 26.914 43.640 27.200 18.46
226	ASP	CA	26.637 42.515 26.346 18.64
227	ASP	С	27.079 41.224 27.054 16.04
228	ASP	0	28.233 41.010 27.385 17.44
229	ASP ASP	CB CG	27.372 42.760 25.036 23.76 26.832 41.802 23.989 28.93
230 231	ASP	OD1	25.701 41.295 24.133 29.41
232	ASP	OD2	27.537 41.564 23.023 33.21
233	PHE	N	26.075 40.401 27.320 14.88
234	PHE	CA	26.312 39.115 27.938 12.14
235 236	PHE	C O	25.696 37.991 27.085 10.36 24.778 38.182 26.301 14.29
237	PHE	СВ	25.708 39.121 29.344 9.41
238	PHE	CG	26.277 40.180 30.227 10.05
239	PHE	CD1	27.508 39.992 30.862 12.58
240	PHE	CD2	25.566 41.344 30.471 6.85
241 242	PHE PHE	CE1 _CE2	28.002 40.930 31.768 9.43 26.045 42.265 31.390 5.14
243	PHE	CZ	27.251 42.063 32.036 7.18
244	PRO	N	26.241 36.762 27.253 7.14
245	PRO	CA	25.755 35.675 26.473 6.52
246 247	PRO PRO	C O	24.277 35.394 26.679 9.69 23.748 35.569 27.762 12.15
4m T F		_	20.7 10 00.000 27.702 12.10

248	PRO	CB	26.607 34.503 26.897 3.58
249	PRO	CG	27.467 34.928 28.059 4.26
250	PRO	CD	27.366 36.422 28.133 3.15
251	CYS	N	23.626 34.982 25.597 12.69
252	CYS	CA	22.261 34.498 25.720 14.65
253	CYS	C	22.172 33.102 25.112 17.19
254	CYS	O	21.342 32.783 24.256 16.47
255	CYS	CB	21.300 35.454 25.016 13.35
256	CYS	SG	21.382 37.237 25.396 15.48
257	ARG	N	23.129 32.258 25.541 18.46
258	ARG	CA	23.174 30.921 24.986 19.34
259	ARG	C	21.865 30.155 25.153 19.02
260	ARG	O	21.360 29.596 24.201 20.59
261	ARG	CB	24.339 30.190 25.590 24.24
262	ARG	CG	25.684 30.614 24.976 34.28
263	ARG	CD	26.506 31.531 25.846 42.36
264	ARG	NE	26.067 31.510 27.243 47.65
265	ARG	CZ	26.832 31.598 28.306 47.20 28.125 31.583 28.117 46.56 26.311 31.717 29.498 44.55
266	ARG	NH1	
267	ARG	NH2	
268	VAL	N	21.246 30.117 26.340 17.94
269	VAL	CA	20.046 29.252 26.364 16.39
270	VAL	C	18.870 29.732 25.445 16.62
271	VAL	O	18.228 28.961 24.769 20.22
272	VAL	CB	19.708 28.709 27.746 14.40
273 274	VAL VAL	CG1 CG2	18.301 29.008 28.203 13.66 20.727 29.076 28.805 11.01 18.666 31.040 25.380 16.61
275 276 277	ALA ALA ALA	N CA C	17.795 31.721 24.438 13.61 18.036 31.352 23.035 12.12
278	ALA	O	17.160 31.403 22.187 12.55 18.080 33.255 24.451 10.37 19.303 31.043 22.828 11.81
279	ALA	CB	
280	LYS	N	
281	LYS	CA	19.667 30.701 21.490 12.07
282	LYS	C	19.712 29.217 21.190 13.74
283	LYS	O	19.996 28.858 20.061 17.30 20.968 31.391 21.154 12.40 20.822 32.884 20.934 13.01
284	LYS	CB	
285	LYS	CG	
286	LYS	CD	19.574 33.099 20.085 17.78
287	LYS	CE	19.498 34.385 19.324 22.14
288	LYS	NZ	18.143 34.532 18.753 25.46
289	LEU	N	19.358 28.374 22.149 13.01
290	LEU	CA	19.324 26.958 21.822 13.50
291	LEU	C	18.159 26.587 20.853 16.14
292	LEU	O	17.057 27.133 20.907 14.24
293 - 294 295	LEU LEU	CB CG CD1	19.105 26.163 23.118 11.77 -20.222 26.230 24.139 8.45 21.462 25.514 23.666 2.00
296 297	PRO	CD2 N	19.752 25.632 25.474 5.67 18.411 25.594 19.966 18.99 17.420 25.201 18.974 20.81
298	PRO	CA	16.038 24.817 19.486 21.23
299	PRO	C	

					321	•
300 301 302 303 304 305 307 308 309 311 312 313 314 315 316 317 318 319 320 321 322 323 324 325 327 328 329 331 331 331 331 331 331 331 331 331 33	LYS LYS ASN ASN ASN ASSN LYS SSN ASN ASN ASN ASN ASN ASN ASN ASN AS	O C C C C C C C N N C C O C C O N N C C O C C C N N C C O C C C O N N C C O C C C N N C C O C C C N N C C O C C C N N N C C O C C C N N N C C O C C C N N N N	15.028 18.053 19.544 19.693 16.032 14.730 13.860 12.648 14.945 15.755 15.710 16.801 16.535 14.556 13.870 14.739 14.821 13.969 14.823 14.213 14.555 15.066 14.971 14.555 15.066 14.971 17.091 19.708 19.204 10.452 10	24.226 24.935 24.245 23.878 25.101 25.010 22.949 23.579 22.689 23.034 22.399 26.233 27.431 28.352 29.444 27.389 26.599 27.678 27.923 28.846 29.452 30.654 29.179 28.542 29.579 29.087 28.544 29.179 28.544 29.179 28.545 29.579 29.666 27.796 26.756 26.857 25.772 29.703 30.375 31.846 32.638 27.622 26.378 25.939 26.631 24.830	18.853 18.312 18.419 19.760 20.703 21.253 21.665 21.812 22.437 23.575 24.822 25.823 27.128 21.22 21.354 23.298 24.621 24.993 25.307 19.857 18.745 18.860 18.781 17.377 16.316 14.949 13.926 12.537 19.113 19.134 20.495 20.711 18.681 19.791 20.899 13.926 12.554 23.827 24.143 25.391 26.964 27.205	22.01 20.31 21.95 20.41 22.87 25.57 19.35 21.88 34.05 18.29 16.34 18.28 15.25 18.39 16.34 19.25 18.39 16.34 17.25 18.35
349	ASN	N	10.877	•		
350	ASN	CA	11.276	33.639	21.283	10.48
351	ASN	<b>C</b> .			20.189	

352	ASN	0	10.425 33.924 19.054 12.34
353	ASN	CB	12.749 33.695 20.923 9.53
354	ASN	CG	13.512 33.223 22.114 13.73
355	ASN	OD1	13.255 33.613 23.245 15.94 14.468 32.378 21.802 13.53
356	ASN	ND2	
·357	ARG	N	9.906 35.498 20.587 9.18
358 359	ARG ARG	CA	9.980 36.947 18.540 6.85
360	ARG	O	9.628 37.017 17.379 6.50
361	ARG	CB	8.232 37.214 20.383 7.45
362	ARG	CG	7.408 38.017 19.428 5.78 6.595 39.001 20.150 7.80
363	ARG	CD	
364	ARG	NE	5.520 38.392 20.894 7.84
365	ARG	CZ	4.356 38.086 20.319 6.71
3 <b>6</b> 6	ARG	NH1	4.093 38.311 19.075 3.66
367	ARG	NH2	3.466 37.541 21.012 3.47
368	TYR	N	11.110 37.522 18.982 8.09
369	TYR	CA	11.994 38.167 18.021 7.20
370	TYR	C	13.398 37.575 18.097 10.87 13.994 37.455 19.164 11.24
371	TYR	O	
372	TYR	CB	12.115 39.628 18.386 5.76
373	TYR	CG	10.820 40.320 18.388 7.25
374	TYR	CD1	10.123 40.388 17.212 10.32
375		CD2	10.304 40.862 19.541 3.38
376	TYR	CE1	8.881 40.994 17.151 12.06
377	. TYR	CE2	9.097 41.526 19.474 8.53
378	∽TYR	CZ	8.354 41.606 18.291 12.52
379	TYR	OH	7.116 42.223 18.267 12.22 13.917 37.304 16.886 12.71
380	ARG	N	
381	ARG	CA	15.268 36.791 16.740 14.19
382	ARG	С	16.325 37.589 17.518 15.21
383	ARG	О	17.239 36.987 18.070 17.69
384	ARG	CB	15.618 36.796 15.242 16.20
385	ARG	CG	16.894 36.009 14.934 20.90
386	ARG	CD	17.513 36.381 13.600 24.13 18.032 37.734 13.679 31.45
387	ARG	NE	
388	ARG	CZ	18.180 38.440 12.559 35.35
389	ARG	NH1	18.031 37.849 11.400 34.27
390	ARG	NH2	18.426 39.722 12.626 36.59
391	ASP	N	16.129 38.922 17.483 15.52
392	ASP	CA	17.068 39.945 17.947 14.40
393	ASP	C	16.939 40.307 19.430 12.23
394	ASP		17.780 41.001 19.966 11.34
395	ASP	СВ	16.825 41.235 17.144 19.70
396	ASP	CG	17.502 41.193 15.756 24.49 18.205 40.237 15.418 27.69
397	ASP	OD1	
398	···-ASP	OD2	17 <del>.</del> 350 42 <del>.</del> 135 14.999 25.18 15.836 39.831 20.061 13.80
399	VAL	N	
400	VAL	CA	15.628 40.144 21.478 13.65
401	VAL	C	15.556 38.911 22.396 13.05
402	VAL	0	14.676 38.066 22.356 10.76
403	VAL	CB	14.512 41.187 21.704 14.37

404 405 406	VAL VAL SER	CG1 CG2 N	13.751 41.144 23.012 10.38 13.935 41.930 20.498 12.19 16.585 38.852 23.257 14.06
407 408 409	SER SER SER	CA C O	16.846 37.705 24.127 9.73 17.249 38.159 25.487 8.28 17.770 39.230 25.677 5.98 17.993 36.869 23.463 10.62
410 411 412	SER SER PRO	CB OG N CA	17.993 36.869 23.463 10.62 17.811 36.472 22.071 11.36 16.972 37.263 26.461 7.35 17.450 37.352 27.792 8.44
413 414 415 416	PRO PRO PRO PRO	CA C O CB	18.908 36.905 27.921 11.00 19.259 35.821 27.503 13.36 16.474 36.351 28.513 7.21
417 418 419	PRO PRO PHE	CG CD N	16.182 35.277 27.531 11.53 16.259 36.046 26.224 9.12 19.753 37.763 28.507 10.36
420 421 422	PHE PHE PHE	CA C	21.035 37.197 28.928 8.56 20.847 35.981 29.853 10.75 19.894 35.899 30.634 12.42
423 424 425	PHE PHE PHE	CB CG CD1	21.829 38.225 29.725 6.92 21.969 39.566 29.090 4.90 22.372 39.651 27.773 2.00
426 427 428	PHE PHE PHE	CD2 CE1 CE2	21.749 40.718 29.841 3.60 22.581 40.893 27.214 3.30 21.964 41.964 29.276 2.19
429 430 431	PHE ASP - ASP	CZ N CA	22.390 42.047 27.962 2.00 21.804 35.056 29.764 11.92 21.710 33.883 30.620 12.44
432 433 434	ASP ASP	C O CB	21.749 34.248 32.129 10.45 21.055 33.664 32.955 14.63 22.852 32.934 30.260 13.15
435 436. 437 438	ASP ASP ASP HIS	CG OD1 OD2 N	22.759 32.466 28.829 17.42 21.740 31.969 28.404 17.02 23.745 32.592 28.162 18.42 22.577 35.242 32.485 6.99
439 440 441	HIS HIS HIS	CA C O	22.781 35.415 33.933 7.65 21.522 35.922 34.692 11.47 21.329 35.728 35.891 12.59
442 443 444	HIS HIS HIS	CB CG ND1	23.994 36.314 34.157 4.46 23.699 37.794 34.137 5.10 23.259 38.489 35.219 8.84
445 446 447	HIS HIS	CD2 CE1 NE2	23.892 38.696 33.091 5.96 23.198 39.775 34.852 5.13 23.568 39.921 33.577 7.85
448 449 450	SER SER SER	N CA C	20.686 36.648 33.929 11.96 19.591 37.347 34.601 10.61 18.221 36.722 34.179 11.50 17.161 37.166 34.622 13.21
451 452 453 454	SER SER SER ARG	O CB OG N	19.578 38.795 34.005 6.54 19.426 38.747 32.546 11.00 18.259 35.718 33.274 10.94
455	ARG	CA	16.999 35.170 32.768 12.70

456	ARG	С	16.139 34.506 33.898 14.39
450 457	ARG	0	16.647 33.926 34.844 12.31
458	ARG	СВ	17.292 34.168 31.637 12.58
459	ARG	CG	17.894 32.846 32.135 14.19
460	ARG	CD	18.073 31.815 31.023 15.27
461	ARG	NE	18.531 30.520 31.560 15.37
462	ARG	CZ	17.720 29.463 31.683 15.74
463	ARG	NH1	16.468 29.566 31.330 13.61
464	ARG	NH2	18.150 28.339 32.167 13.65
465	ILE	N	14.811 34.584 33.745 16.19
466	ILE	CA	13.964 33.837 34.671 13.97
467	ILE	С	13.863 32.356 34.253 13.70
468	ILE	0	13.599 32.046 33.119 15.22
469	ILE	CB	12.583 34.496 34.725 14.29
470	ILE	CG1	12.697 35.981 35.109 14.68
471	ILE	CG2	11.695 33.761 35.745 14.23 12.768 36.193 36.625 11.53
472 473	LYS	CD1 N	14.079 31.452 35.184 15.41
474	LYS	CA	13.901 30.039 34.865 16.79
475	LYS	C	12.555 29.531 35.391 17.35
476	LYS	Ö	12.222 29.772 36.539 19.31
477	LYS	СВ	14.952 29.273 35.664 17.97
478	LYS	CG	16.348 29.641 35.224 20.00
479	LYS	CD	17.342 28.756 35.923 21.07
480	LYS	CE	18.757 29.183 35.586 25.86
481	LYS	NZ	19.661 28.132 36.065 29.41
482	LEU	N	11.827 28.807 34.546 17.14
483	LEU	CA	10.659 28.096 35.026 16.17 11.101 26.982 35.997 18.46
484 485	LEU LEU	C O	11.101 26.982 35.997 18.46 12.139 26.361 35.815 16.31
486	LEU	CB	9.925 27.512 33.820 16.73
487	LEU	CG	9.241 28.474 32.838 16.01
488	LEU	CD1	9.233 28.140 31.340 18.22
489	LEU	CD2	9.174 29.942 33.208 19.99
490	HIS	N	10.307 26.720 37.031 20.86
491	HIS	CA	10.504 25.539 37.856 22.12
492	HIS	С	9.967 24.256 37.160 26.50
493	HIS	0	9.214 23.488 37.729 29.06
494	HIS	CB	9.795 25.710 39.208 20.80
495	HIS	CG ND1	10.244 26.940 39.961 18.49 9.650 27.355 41.096 18.74
496 497	HIS HIS	CD2	11.279 27.836 39.657 17.16
498	HIS	CE1	10.306 28.462 41.471 16.79
499	HIS	NE2	11.293 28.776 40.626 15.80
500	GLN	N	10.416 24.000 35.928 31.86
501	GLN	CA	9.989 22.769 35.260 36.18
502	GLN	uC ,	11.226_22.046_34.724_37.67
503	GLN	0	12.153 22.676 34.251 38.03
504	GLN	СВ	8.969 23.068 34.141 38.37
505	GLN	CG	9.413 24.118 33.110 45.32
506 507	GLN	CD OE1	8.538 24.115 31.844 51.67 8.913 23.628 30.781 53.74
507	GLN	OE1	8.913 23.628 30.781 53.74

508 509 510 511 512 513 514 515 516 517 518 519 520 521 522 523 524 525 526	GLN GLU GLU GLU GLU GLU GLU ASP ASP ASP ASP ASP ASP ASP ASP	NE2 N CA C O CB CCD OE1 OE2 N CA C O CB CG OD1 OD2 N	7.325 24.611 32.016 51.04 11.195 20.711 34.825 40.34 12.374 19.917 34.434 41.41 12.545 19.806 32.896 40.03 13.654 19.763 32.381 38.15 12.223 18.535 35.054 45.95 12.189 18.568 36.590 52.82 11.488 17.315 37.138 60.20 11.234 16.382 36.370 63.99 11.181 17.291 38.322 61.75 11.382 19.779 32.207 39.83 11.403 19.725 30.735 37.84 12.238 20.863 30.075 31.75 13.278 20.692 29.439 32.35 9.941 19.774 30.258 45.12 9.937 19.953 28.726 53.29 10.589 19.165 28.040 57.18 9.363 20.939 28.246 56.58 11.695 22.072 30.254 26.65
527 528 529 530 531 532 533 534 535 536 537 538 539 540 541 542 543 544 545 546 547 548 549 550 551 552 553	ASN ASN ASN ASN ASN ASP ASP ASP ASP ASP ASP TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	CA C O CB CG OD1 N CA C O CB CG OD2 N CA C O CB CD1 CD2 CE1 CE2 CZ OH N	12.390 23.201 29.678 21.86 12.099 24.449 30.528 20.05 11.018 25.025 30.494 21.82 11.951 23.347 28.213 18.23 12.778 24.425 27.560 15.37 13.328 25.282 28.216 19.18 12.889 24.375 26.266 5.53 13.138 24.890 31.233 17.24 12.969 26.055 32.084 15.50 12.983 27.419 31.339 13.37 12.998 28.461 31.970 16.65 14.049 26.012 33.173 19.65 15.459 26.514 32.718 24.75 15.644 27.054 31.628 25.78 16.425 26.355 33.444 27.00 13.057 27.417 29.999 11.69 13.282 28.673 29.271 9.17 11.981 29.477 29.069 8.95 10.979 28.944 28.618 10.59 14.044 28.417 27.937 9.39 14.285 29.690 27.182 7.68 15.282 30.541 27.605 5.47 13.461 30.068 26.121 6.70 15.383 31.821 27.085 6.80 13.539 31.330 25.557 5.20 14.498 32.243 26.073 8.53 14.596 33.555 25.625 8.14 12.050 30.770 29.393 8.87
555 556 557 558	ILE ILE ILE ILE	CA C O CB	11.093 31.830 29.070 10.50 11.899 33.107 28.707 10.84 12.881 33.469 29.342 11.65 10.060 32.108 30.209 11.32

611 612 613 614 615 616 617 619 619 621 621 621 622 623 624 625 626 627 632 633 635 636 637 638 639 640 641 642 643 644 645 646 647 648 649 649 649 649 649 649 649 649 649 649	MET TUUUUUUUUUUUUUUUUUUUU AAAAAANNNNNNNNNN	CSCNCCOCCOONCCOCCOONCCOCNCCOCCONNCCOCCONNCCOCCO	16.626 48.573 39.119 13.99 18.121 48.583 38.091 17.00 18.404 50.363 38.012 4.51 19.696 47.328 42.741 19.80 20.201 47.101 44.094 21.74 20.060 48.353 44.993 21.70 19.551 48.311 46.106 21.22 21.652 46.627 44.007 24.29 22.273 46.291 45.379 31.56 23.780 45.945 45.252 36.04 24.575 46.709 44.678 39.83 24.167 44.887 45.726 40.03 20.500 49.491 44.427 22.15 20.486 50.733 45.200 22.50 19.056 51.237 45.483 22.61 18.690 51.633 46.586 23.83 21.321 51.751 44.448 23.01 21.465 53.095 45.163 29.10 21.921 54.170 44.157 35.50 21.836 53.969 42.942 36.25 22.346 55.218 44.595 38.56 18.218 51.181 44.432 23.89 16.788 51.506 44.582 21.87 16.051 50.494 45.447 20.77 15.078 50.857 46.077 21.59 16.112 51.586 43.207 19.61 16.555 49.262 45.482 22.52 15.859 48.190 46.212 23.75 14.447 47.831 45.617 21.92 13.548 47.415 46.324 23.13 15.781 48.573 47.700 28.00 17.090 48.375 48.482 37.73 17.102 46.993 49.161 45.98 16.962 46.865 50.359 49.70 17.222 45.958 48.347 46.64 14.293 47.996 44.288 19.87 13.167 47.433 43.520 15.56 13.671 46.530 42.369 15.93 14.589 46.892 41.649 19.66 12.378 48.566 42.894 12.31 11.026 48.114 42.359 8.59 9.975 49.222 42.284 8.75 9.454 49.581 43.583 9.44
649	ARG	O	14.589 46.892 41.649 19.66
650	ARG	CB	12.378 48.566 42.894 12.31
652	ARG	CD	9.975 49.222 42.284 8.75
654	ARG	CZ	8.781 50.712 43.804 13.29
655	ARG	NH1	8.507 51.573 42.868 14.59
656	ARG	NH2	8.376 50.970 44.997 14.52
657	SER	N	13.018 45.374 42.191 13.03
658	SER	CA	13.174 44.563 40.969 11.10 11.905 44.484 40.133 9.85
659	SER	С	10.797 44.430 40.625 7.17
660	SER	О	
661	SER	CB	13.491 43.098 41.352 9.97
662	SER	OG	14.257 42.832 42.565 16.33

663 664 665 666 667 668 669 670 671 672 673 674 675 676 677 678 679	TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	N CA C O CB CD1 CD2 CE1 CE2 CZ OH N CA C O CB	12.149 44.304 38.838 10.99 11.059 43.986 37.924 11.25 11.428 42.738 37.134 11.69 12.600 42.426 36.947 12.93 10.855 45.145 36.944 11.85 11.048 46.504 37.543 11.61 12.330 46.990 37.697 13.15 9.974 47.284 37.928 10.09 12.574 48.223 38.239 12.30 10.189 48.561 38.453 11.52 11.477 49.041 38.615 11.57 11.518 50.318 39.128 16.85 10.411 42.065 36.626 10.13 10.683 41.103 35.561 10.15 10.210 41.646 34.208 11.23 9.017 41.882 34.001 14.10 9.984 39.783 35.918 8.65
680 681 682 683 684 685 686 687 688 689 691 692 693 694 695 696 697 701 702 703 704 705 707 708 709 711 712 713	ILE LEUU UUU UU RRRRRRRRNNNNNNNNNNNNNNNNNNNNN	CG1 CG2 CD1 N CC O CBG1 CC O CBG1 CC O CBG CC O N CC O CBG CC O N	10.575       39.226       37.221       5.54         10.117       38.788       34.758       4.72         9.885       37.981       37.750       3.41         11.171       41.837       33.292       8.83         10.757       42.279       31.952       8.27         10.642       41.089       31.042       9.63         11.563       40.274       30.925       10.20         11.754       43.266       31.361       6.64         11.554       44.667       31.933       5.33         12.441       45.790       31.363       7.20         11.164       44.814       33.411       7.37         9.476       41.020       30.372       9.19         9.353       39.941       29.424       7.94         8.748       40.395       28.068       7.09         8.236       41.487       27.941       6.68         8.633       38.725       30.140       7.51         8.460       37.446       29.475       10.84         7.202       39.983       25.785       4.05         6.657       39.538       25.856       8.34         6.182       38.810

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714	PRO	0	2.432	36.210 23.249 9.13
715			2.432	35.533 26.387 7.90
	PRO	CB		
716	PRO	CG	2.452	36.716 27.320 8.31
717	PRO	CD '	3.076	37.939 26.655 8.22
718	LEU	N	3.826	34.451 23.462 11.50
719	LEU	CA	3.453	33.772 22.222 12.63
720	LEU	C .	2.284	32.817 22.537 10.92
721	LEU	0	2.099	32.425 23.675 10.18
722	LEU.	CB	4.653	32.966 21.630 12.16
723	LEU	CG	5.658	33.617 20.677 10.05
724	LEU	CD1	7.130	33.604 21.085 9.84
725	LEU	CD2	5.220	34.724 19.751 8.39
726	PRO	N	1.493	32.446 21.503 11.77
727	PRO	CA	0.410	31.494 21.723 12.03
	PRO	C	0.830	30.158 22.433 13.54
728				
729	PRO	0	0.089	
730	PRO	СВ	-0.132	31.242 20.320 9.15
731	PRO	CG	0.277	32.454 19.478 10.31
732	PRO	CD	1.572	32.919 20.123 10.84
733	ASN	Ν	2.100	29.779 22.259 13.43
734	ASN	CA	2.585	28.584 22.964 13.13
735	ASN	C .	3.324	28.859 24.291 11.81
736	ASN	0	3.962	27.974 24.845 11.04
737	-ASN	CB	3.496	27.799 22.046 13.99
738	ASN	CG	4.657	28.667 21.597 16.96
739	ASN	OD1	4.503	29.783 21.131 22.38
740	ASN	ND2	5.839	28.120 21.783 18.79
741	THR	N	3.286	30.095 24.782 9.69
742	THR	CA	4.037	30.277 26.042 9.51
743	THR	C	3.210	31.147 27.020 11.20
744	THR	Ö.	3.752	31.735 27.945 11.31
745	THR	СВ	5.272	31.194 25.696 8.41
746	THR	OG1	4.946	32.545 25.282 10.58
		CG2	6.132	30.535 24.591 5.31
747	THR			31.177 26.815 12.27
748	CYS	N	1.877	31.799 27.822 12.45
749	CYS	ÇA	1.007	
750	CYS	С	0.947	30.973 29.109 12.47
751	CYS	0	0.771	31.526 30.183 14.18
752	CYS	CB	-0.422	32.004 27.341 8.19
753	CYS	SG	-0.524	32.997 25.849 9.18
754	GLY	N	1.156	29.654 28.975 10.77
755	GLY	CA	1.293	28.814 30.180 10.51
756	GLY	С	2.589	29.054 30.979 13.69
757	GLY	Ο.	2.584	29.199 32.182 14.84
758	HIS.	Ν	3.719	29.198 30.259 14.08
759	HIS	CA	4.993	29.584 30.868 11.53
760	HIS	С	4.914	30.988 31.502 11.94
761	HIS	0	5.482	31.244 32.554 14.08
762	HIS	СВ	6.065	29.732 29.789 10.46
763	HIS	CG	6.166	28.534 28.894 10.11
764	HIS	ND1	6.494	28.651 27.599 10.39
765	HIS	CD2	5.952	27.176 29.154 10.94
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818	TRP	CA	2.909	31.996	40.053	12.02
819	TRP	C	4.227	31.749	40.844	11.70
820	TRP	0	4.326	32.034	42.034	12.54
821	TRP	CB	2.217	30.672	39.710	10.42
822	TRP	CG	1.863	29.951	41.001	12.86
823	TRP	CD1	2.696	29.101	41.739	14.68
824	TRP	CD2	0.644	30.071	41.772	14.87
		NE1	2.072	28.715	42.886	15.11
825	TRP					
826	TRP	CE2	0.810	29.275	42.935	15.72
827	TRP	CE3	-0.539	30.710	41.553	15.15
828	TRP	CZ2	-0.194	29.217	43.858	16.05
829	TRP	CZ3	-1.558	30.644	42.477	14.60
830	TRP	CH2	-1.389	29.889	43.623	17.06
831	GLU	N	5.226	31.223	40.140	12.34
832	GLU	CA	6.504	30.890	40.783	13.26
833	GLU	C	7.246	32.102		15.44
834	GLU	0	7.808	32.021	42.444	18.46
835	GLU	СВ	7.401	30.155	39.785	11.02
836	GLU	CG	6.906	28.742	39.525	9.86
837	GLU	CD	7.513	28.097	38.292	10.94
838	GLU	OE1	8.355	28.663	37.634	13.74
839	GLU	OE2	7.119	26.985	38.017	12.82
840	GLN	N	7.186		40.614	15.26
841	GLN	CA	7.871	34.449		15.75
842	GLN	C	7.052	35.265	42.051	16.17
843	GLN	0	7.555	36.199	42.665	16.73
844	GLN	CB	8.228	35.289	39.806	16.31
845	GLN	CG	9.155	34.556	38.822	16.92
846 '	GLN	CD .	10.425	34.083	39.552	19.20
847	GLN	OE1	11.001	34.783	40.362	19.09
848	GLN .	NE2	10.810	32.870	39.299	15.74
849	LYS	N	5.771	34.861	42.227	15.25
850	LYS	CA	4.934	35.484	43.254	13.27
851	LYS	C	4.651	36.977		13.03
852	LYS	0	4.524	37.792		10.75
853	LYS	СВ	5.567	35.249		16.84
854		,	5.787		44.877	20.72
	LYS	CG				
855	LYS	CD	6:097	33.446		23.31
856	LYS	CE	6.774		46.522	
857	LYS	·NZ	7.947		45.619	
858	SER	N	4.552	37.327	41.701	14.11
859	SER	CA	4.152	38.693	41.351	13.84
860	SER	С	2.681	38.899	41.621	14.27
861	SER	Ō	1.854		41.464	13.39
862	SER	СВ	4.462		39.829	13.28
863	SER	OG	5.745	38.417		13.97
864	ARG	N	2.389	40.117		15.30
865	ARG	CA	1.022			15.25
866	ARG	C	0.390		41.189	14.31
867	ARG	0	-0.817	41.375		11.90
868	ARG	CB	1.055		43.620	16.01
869	ARG	CG	-0.333	41.453	44.243	21.04

		0.5	0.004	40 000 40 000 00 74	
870 871	ARG ARG	CD NE		42.809 43.980 23.71 43.095 44.935 25.68	
872	ARG	CZ		42.448 44.880 25.46	
873	ARG	NH1		41.374 44.164 22.79	
874	ARG	NH2		42.930 45.529 27.74	
875	GLY	N	1.276	41.733 40.301 12.62	
876	GLY	CA		42.673 39.263 10.16	
877	GLY	C ·		42.376 37.956 10.24	
878	GLY	0		41.984 37.944 9.74	
879	VAL	N	0.909 1.554	<b>42</b> .597 36.861 10.10 <b>42</b> .588 35.545 9.04	
880 881	VAL VAL	CA C	1.189	43:904 34.860 7.92	
882	VAL	0	0.021	44.276 34.795 8.52	
883	VAL	СВ	0.955	41.434 34.717 7.21	
884	VAL	CG1	0.741	40.059 35.377 10.00	
885	VAL	CG2	1.208	41.428 33.216 6.46	
886	VAL	N	2.225	44.560 34.356 6.18	
887	VAL	CA	2.002	45.766 33.558 5.67	
888	VAL VAL	C O	2.204 3.294	45.440 32.090 5.05 45.104 31.685 8.26	
889 890	VAL	CB	2.942	46.867 34.051 4.27	
891	VAL	CG1	2.627	47.146 35.522 5.59	
892	VAL	CG2	2.821	48.193 33.311 5.52	
893	MET	N	1.129	45.564 31.315 6.05	
894	MET	CA	1.132	45.435 29.850 6.12	
895 896	MET MET	C O	1.159 0.263	46.791 29.168 5.90 47.591 29.379 6.94	
897	MET	CB	-0.222	44.895 29.428 6.49	
898	MET	CG	-0.347	44.620 27.938 5.59	
899	MET	SD	-1.592	43.356 27.620 10.50	
900	MET	CE	-1.470	43.249 25.807 2.00	
901	LEU	N	2.146	47.028 28.319 5.40	
902 903	LEU	CA	2.242 1.866	48.428 27.799 5.78 48.558 26.299 7.49	
903	LEU	Ö	2.091	49.581 25.666 7.99	
905	LEU	СВ	3.632	48.997 28.075 6.33	
906	LEU	CG	3.952	49.053 29.593 8.42	
907	LEU	CD1	3.064	50.026 30.390 6.57	
908	LEU ASN	CD2 N	5.400 1.336	49.460 29.834 5.25 47.445 25.747 6.12	
909 910	ASN	CA	1.077	47.367 24.327 8.78	
911	ASN	C	-0.348	46.840 24.026 10.77	
912	ASN	O .	-0.979		
913	ASN	СВ	2.125	46.458 23.685 10.20	
914	ASN	CG	1.956	45.019 24.175 10.50	
915 916	ASN -ASN	OD1 ND2	1.469 2.370	44.137 23.509 13.52 44.830 25.401 11.44	
917	ARG	NDZ		46.974 22.777 12.13	
918	ARG	CA	-1.993		
919	ARG	C	-1.619	44.915 21.703 11.64	ļ
920	ARG	0		44.743 21.149 13.53	
921	ARG	СВ	-2.716	47.222 21.340 15.99	)

922 923 925 927 929 931 932 933 935 937 939 939 941 942 943 945 949 951 953 954 955 957 958 968 968 969 968	ARG ARG ARG VALL VALL TTTTTTTTUUUUUUUUUUSSSSSSSSSSSSSSSSSSS	CCNCNNNCCOCCONCCOCCSCNCCOCCCONCCOCCCNNCCONCCO	-4.252 -4.304 -3.229 -2.041 -3.324 -2.522 -2.168 -2.076 -1.410 -3.314 -3.544 -3.739 -2.781 -2.503 -1.936 -2.532 -3.734 -3.319 -4.505 -4.816 -0.776 -0.179 0.459 1.028 0.888 0.407 1.416 2.568 1.021 0.322 0.947 0.631 1.464 2.450 2.681 3.794 4.085 2.832 -0.656 -1.177 -0.875 -1.358 -0.076 0.409 0.010 -0.199 1.956	49.313 49.380 49.394 49.594 49.596 49.6400 49.6400 49.6	19.246 50.26 19.777 51.39 17.948 53.49 21.780 11.90 21.057 13.18 19.536 12.04 18.850 12.44 21.330 14.90 22.786 14.45 20.196 13.17 19.030 13.86 17.669 16.37 17.600 15.93 18.063 15.73 16.776 19.13 15.289 25.05 14.175 30.99 13.287 28.20 16.938 15.17 16.644 16.14 15.260 18.85 14.830 20.26 17.679 15.90 19.109 17.76 20.137 18.82 19.990 16.03 21.072 15.57 14.571 22.41 13.227 23.76 12.303 21.76 11.599 21.77 13.386 24.05 13.822 23.31 14.847 25.30 15.113 28.40 15.212 27.38 12.413 20.87 11.522 19.66 11.919 19.12 11.269 19.90 12.978 17.43 13.385 16.48 14.786 15.10 15.622 15.11 13.330 18.82
966 967 968	SER SER SER	CA C O	0.409 0.010 -0.199 1.956 2.476 -0.028 -0.222	42.896 42.558 43.426 42.955	13.385 16.48 14.786 15.10 15.622 15.11 13.330 18.82 12.036 30.66 14.996 14.24 16.300 13.02

074		0	2.026 39.930 16.888 12.03
974 975	LEU LEU	СВ	-0.717 39.195 16.092 13.63
976	LEU	CG	-2.160 39.215 15.545 10.24
977	LEU	CD1	-3.226 40.017 16.301 10.97
978	LEU LYS	CD2 N	-2.646 37.927 14.916 9.61 1.105 41.571 18.084 11.18
979 980	LYS	CA	2.290 41.882 18.858 8.96
981	LYS	C	2.390 41.095 20.161 7.30
982	LYS	0	3.376 41.101 20.884 6.49
983	LYS LYS	CB CG	2.258 43.376 19.126 12.36 2.527 44.152 17.844 14.52
984 985	LYS	CD	3.978 44.105 17.484 16.12
986	LYS	CE	4.345 44.924 16.271 20.42
987	LYS	NZ	5.804 45.081 16.202 23.02
988	CYS	N .	1.319 40.356 20.402 6.54 1.256 39.673 21.686 10.66
989 990	CYS CYS	CA C	0.100 38.668 21.635 14.07
991	CYS	Ö	-0.882 38.889 20.936 18.36
992	CYS	СВ	1.133 40.710 22.839 9.20
993	CYS ALA	SG N	0.873 40.094 24.504 5.00 0.214 37.548 22.367 13.71
994 995	ALA	CA	-0.970 36.706 22.498 11.28
996	ALA	C	-2.054 37.332 23.451 13.57
997	ALA	0	-1.787 38.061 24.405 13.87
9 <b>98</b> 9 <b>9</b> 9	ALA GLN	CB N	-0.506 35.347 22.988 5.96 -3.315 36.986 23.148 13.22
1000	GLN	CA	-4.355 37.267 24.134 11.86
1001	GLN	·Č · ···	-4.194 36.315 25.338 9.90
1002	GLN	O CB	-4.748 35.227 25.329 8.51 -5.696 36.958 23.442 11.64
1003 1004	GLN GLN	CG	-6.901 37.307 24.331 12.96
1005	GLN	CD	-6.930 38.782 24.786 15.38
1006	GLN	OE1	-7.275 39.155 25.885 18.48
1007	GLN TYR	NE2- N	-6.501 39.645 23.923 10.76 -3.353 36.690 26.311 10.20
1008 1009	TYR	CA	-2.959 35.652 27.305 11.34
1010	TYR	С	-3.874 35.622 28.587 10.00
1011	TYR	0	-3.692 34.809 29.480 12.22
1012 1013	TYR TYR	CB CG	-1.451 35.764 27.671 9.57 -1.092 37.093 28.295 10.61
1014	TYR	CD1	-1.112 37.261 29.674 11.35
1015	TYR	CD2	-0.748 38.189 27.492 10.24
1016	TYR TYR	CE1 CE2	-0.865 38.489 30.277 10.01 -0.500 39.429 28.066 8.22
1017 1018	TYR	CZ	-0.564 39.588 29.448 8.65
1019	TYR	ОН	-0.317 40.841 29.935 7.28
	TRP	N	-4.815-36.557-28.642-9.43 5.744-36.600-30.703-10.16
1021 1022	TRP TRP	CA C	-5.714 36.690 29.792 10.16 -7.175 36.690 29.287 10.75
1023	TRP	Ö	-7.442 37.154 28.187 8.67
1024	TRP	СВ	-5.341 37.945 30.612 8.10
1025	TRP	CG	-5.788 39.222 29.929 8.50

1026 1027 1028 1029 1030 1031 1032 1033 1034 1035 1036 1037 1038 1040 1041 1042 1043 1044 1045 1046 1047 1050 1051 1052 1053 1054 1055 1056 1067 1068	TRP P P P P P P O O O O O O O O O O O O O	CD1 CD2 NE2 CZ3 CN CC O CC CN CC O CC CONN CC O CC CNN CC O CC CONN CC CONN CC O CC CONN CC CONN CC O CC CONN CC CON	-6.970 39.931 30.199 9.41 -5.138 39.917 28.835 8.04 -7.089 40.994 29.356 8.02 -5.979 41.032 28.508 8.86 -3.990 39.680 28.138 6.23 -5.626 41.863 27.469 7.39 -3.641 40.518 27.068 8.40 -4.457 41.614 26.747 8.65 -8.137 36.145 30.127 11.94 -9.538 36.107 29.746 11.58 10.158 37.512 29.637 14.25 10.022 38.396 30.486 14.80 10.225 35.238 30.819 8.41 -9.311 35.278 32.027 5.08 -7.939 35.615 31.474 10.33 10.892 37.626 28.524 15.45 11.657 38.851 28.328 18.38 13.068 38.858 28.949 19.02 13.747 39.861 28.892 20.54 11.682 39.166 26.841 20.69 10.255 39.453 26.327 27.58 10.336 39.675 24.835 28.97 10.447 38.767 24.036 27.29 10.375 40.949 24.506 28.60 13.516 37.738 29.523 19.10 14.764 37.856 30.269 18.50 14.946 36.717 31.217 15.91 14.342 35.680 31.035 16.84 15.921 37.842 29.309 24.10 15.932 36.706 28.297 25.36 16.993 36.968 27.238 29.33 17.243 38.464 27.014 37.08 18.344 38.699 26.095 42.94 15.801 36.951 32.206 14.50 15.976 36.171 33.393 14.90 16.338 34.709 33.088 18.20 15.719 33.790 33.632 21.55 17.056 36.856 34.217 15.32 16.575 38.151 34.874 13.30 16.724 39.424 34.004 16.29 16.616 39.376 32.782 16.80 16.964 40.491 34.568 16.60 17.317 34.475 32.183 19.29
1063	GLU	CB	17.056 36.856 34.217 15.32
1064	GLU	CG	16.575 38.151 34.874 13.30
1065	GLU	CD	16.724 39.424 34.004 16.29
	GLU	OE2	16.964 40.491 34.568 16.60 17.317 34.475 32.183 19.29 17.820 33.133 31.855 19.57
1070	GLU	C	16.760 32.233 31.213 21.30
1071	GLU	O	16.888 31.007 31.232 20.15
1072	GLU	.CB	18.964 33.205 30.827 18.63
1073	GLU	CG	19.852 34.414 31.027 16.44
1074	GLU	CD	19.393 35.654 30.276 16.06
1075	GLU	OE1	19.651 35.750 29.079 17.52
1076	GLU	OE2	18.763 36.522 30.864 15.56
1077	LYS	N	15.763 32.878 30.564 21.95

4070			
1078	LYS	CA	14.806 32.104 29.783 24.05
1079	LYS	С	13.376 32.240 30.376 24.41
1080	LYS	0	12.539 32.989 29.869 24.56
1081	LYS	CB	14.884 32.562 28.310 26.00
1082.	LYS	CG	16.297 32.545 27.708 32.43
1083	LYS	CD	16.445 31.919 26.320 39.12
1084	LYS	CE	15.519 30.736 26.063 47.59
1085	LYS	NZ	16.271 29.672 25.392 51.15
	GLU		
1086		N	13.140 31.494 31.483 24.94
1087	GLU	CA	11.782 31.372 32.025 23.85
1088	GLU	С	10.831 30.682 31.032 20.25
1089	GLU	0	11.227 29.985 30.119 19.30
1090	GLU	CB	11.754 30.531 33.308 26.32
1091	GLU	CG	13.084 30.345 34.016 32.27
1092	GLU	CD	13.812 29.168 33.377 33.00
1093	GLU	OE1	13.258 28.077 33.359 32.06
1094	GLU	OE2	14.929 29.369 32.909 35.48
1095	MET	N.	-9.550 30.894 31.280 17.49
1096	MET	CA	-8.526 30.188 30.550 15.38
1097	MET	C.	-7.902 29.097 31.442 16.86
1098	MET	0	
			-7.471 29.361 32.554 18.95 -7.480 24.220 20.446 42.45
1099	MET	CB	-7.489 31.239 30.146 12.45
1100	MET	CG	-8.055 32.218, 29.122 11.06
1101	MET	SD	-6.888 33.465 28.615 15.30
1102	MET	CE	-5.835 32.442 27.602 13.48
1103	ILE	N	-7.819 27.879 30.942 15.12
1104	ILE	CA	-6.944 26.908 31.605 16.59
1105	ILE	C	-5.708 26.614 30.770 17.15
1106	ILE	0	-5.788 26.242 29.614 19.86
1107	ILE	CB	-7.708 25.631 31.971 18.08
1108	ILE	CG1	-8.632 25.919 33.163 23.46
1109	ILE	CG2	-6.764 24.493 32.352 17.34
1110	ILE	CD1	10.089 25.551 32.914 24.91
1111	PHE	N.	-4.558 26.751 31.426 16.66
1112	PHE	CA	-3.307 26.461 30.755 16.55
1113	PHE	C	-2.838 25.024 31.123 19.38
1114	PHE	0	•
1115	PHE	·CB	-2.336 27.588 31.105 12.36
1116	PHE	CG	-2.824 28.944 30.818 10.57
1117	PHE	CD1	-2.749 29.439 29.534 8.04
1118	PHE	CD2	-3.334 29.723 31.855 11.55
1119	PHE	CE1	-3.201 30.723 29.284 10.42
1120	PHE	CE2	-3.787 31.007 31.608 11.87
1121	PHE	CZ	-3.719 31.506 30.311 12.28
1122	GLU	N	-3.116 24.104 30.205 21.82
1123	GLU	CA	-3.039 22.720 30.661 25.19
1124	GLU	C	-1.584_ 22.256_30.878_24.14_
1125	GLU	Ō	-1.290 21.361 31.659 25.33
1126	GLU	СВ	-3.639 21.787 29.605 31.07
1127	GLU	CG	-5.166 21.765 29.396 39.94
1128	GLU	CD	-5.454 20.639 28.364 48.24
1129	GLU	OE1	-5.364 20.905 27.164 52.05
20		<u> </u>	5.55 £5.555 £1.104 JZ.05

1130 1131 1132 1133 1134 1135 1136 1137 1138 1139 1140 1141 1143 1144 1145 1151 1152 1153 1154 1163 1164 1165 1167 1168 1170 1171 1172 1173 1174 1175	G A S A S A S A S T T T T T T T T A A A A	OE2 NCCOCCOONCCOCCOCNCCOCCONNCCCOCCONCCCOCCNNCCOCCO	-5.724 19.507 28.772 51.99 -0.697 22.914 30.136 23.00 0.712 22.482 30.155 22.21 1.446 22.782 31.508 20.47 2.261 22.031 32.033 19.77 1.332 23.130 28.915 21.00 1.430 24.643 29.079 22.61 0.486 25.282 29.556 26.49 2.454 25.190 28.734 21.60 1.068 23.914 32.079 19.14 1.645 24.175 33.404 16.22 0.561 24.054 34.479 18.50 0.848 24.327 35.628 20.38 2.051 25.710 33.412 14.66 0.989 26.641 33.065 13.92 3.261 25.983 32.480 11.30 -0.686 23.637 34.105 19.33 -1.730 23.331 35.094 19.95 -2.154 24.539 35.987 21.85 -2.161 24.482 37.214 22.33 -1.180 22.269 36.021 24.72 -2.334 21.494 36.646 27.53 -3.342 21.190 36.060 30.68 -2.193 21.200 37.881 26.04 -2.484 25.635 35.297 22.78 -2.886 26.899 35.936 22.24 -4.265 27.315 35.376 22.59 -4.561 27.167 34.194 23.95 -1.875 28.025 35.599 21.01 -0.701 28.404 36.548 17.04 0.610 28.303 35.804 13.07 -0.612 27.764 37.930 14.04 -5.087 27.901 36.222 20.53 -6.322 28.485 35.725 17.91 -6.339 30.003 35.942 17.68 -5.903 30.515 36.958 16.29 -7.441 27.691 36.368 17.25 -8.807 28.050 35.869 18.58 -9.895 27.178 36.443 22.19 11.140 27.885 36.932 22.34 11.879 26.984 37.836 24.46 -6.846 30.705 34.934 17.39 -7.006 32.157 34.943 15.35 -8.477 32.621 34.637 16.31 -9.075 32.259 33.636 16.43 -5.961 32.752 34.009 12.86
1175 1176 1177 1178 1179	LEU LEU LEU THR	CB CG CD1 CD2 N	-5.961 32.752 34.009 12.86 -5.853 34.284 34.126 9.27 -5.021 34.787 32.956 8.45 -5.200 34.687 35.454 9.53 -9.012 33.479 35.526 13.25
1180 1181	THR THR	CA C	10.363 34.063 35.463 12.10 10.355 35.563 35.597 13.46

1182 1183 1184 1185 1186 1187 1188 1190 1191 1193 1194 1195 1196 1197 1198 1190 1201 1202 1203 1204 1205 1207 1208 1210 1211 1212 1213 1214 1215 1216 1217 1218 1219 1220 1221 1222 1223 1224 1225 1228 1229	TTTTLLLLLLLLLLLLLLLLLLLLLLLLLLLLLLLLLL	O C G G C C C C C C C C C C C C C C C C	-9.718 36.088 36.499 15.11 11.097 33.461 36.694 12.89 10.946 32.013 36.745 11.16 12.589 33.673 36.530 13.73 11.097 36.253 34.707 12.69 11.361 37.680 34.900 13.21 12.268 37.840 36.109 16.36 13.383 37.337 36.138 17.33 12.105 38.280 33.697 9.27 12.355 39.790 33.774 5.96 13.113 40.189 32.520 3.05 11.034 40.574 33.866 2.04 11.767 38.531 37.113 16.98 12.711 38.814 38.209 16.56 13.488 40.095 37.970 15.67 14.677 40.183 38.224 16.31 12.058 38.673 39.567 14.46 11.521 37.227 39.718 13.21 13.057 39.124 40.658 11.57 12.570 36.104 39.539 7.88 12.769 41.053 37.410 16.03 13.492 42.181 36.859 17.37 12.449 43.133 36.298 19.23 11.276 42.946 36.557 19.21 14.107 42.990 38.027 20.04 13.123 43.511 39.003 20.18 12.886 44.193 35.634 22.29 11.982 45.230 35.140 23.93 12.557 46.663 35.235 23.31 13.754 46.861 35.283 27.00 11.701 44.924 33.697 24.38 12.961 44.717 32.896 25.56 12.581 44.588 31.426 31.45 11.402 44.721 31.054 32.20 13.483 44.345 30.651 36.07 11.671 47.634 35.279 21.18 11.995 49.030 35.441 17.82 11.395 49.764 34.225 18.72 10.200 50.051 34.108 20.93 11.443 49.430 36.814 22.34 11.406 50.932 37.034 28.58 12.354 51.602 36.636 27.70 10.412 51.436 37.597 33.90 12.306 49.980 33.282 18.31 11.933 50.554 32.008 19.98 11.874 52.099 32.025 20.84 12.850_52.781_32.267_20.15 12.946 50.053 30.978 20.01
1227	ILE	C	11.874 52.099 32.025 20.84
1228	ILE	O	12.850 52.781 32.267 20.15
1229	ILE	CB	12.946 50.053 30.978 20.01
1230	ILE	CG1	12.853 48.540 30.856 20.02
1231	ILE	CG2	12.762 50.696 29.607 19.35
1232	ILE	CD1	13.722 47.987 29.733 21.08
1233	LYS	N	10.700 52.617 31.682 22.09

1255         TYR         CD1         -5.465         57.864         30.081         14.09           1256         TYR         CD2         -3.682         56.427         29.721         13.30           1257         TYR         CE1         -5.016         58.191         31.311         17.45           1258         TYR         CE2         -3.090         56.866         30.874         19.22           1259         TYR         CZ         -3.767         57.715         31.696         19.65           1260         TYR         OH         -3.079         58.078         32.801         20.24           1261         TYR         N         -6.461         53.351         29.591         15.06           1262         TYR         CA         -5.915         52.151         30.228         14.61           1263         TYR         C         -7.010         51.388         30.962         15.66           1264         TYR         O         -8.023         51.956         31.320         17.43           1265         TYR         CB         -4.719         52.522         31.134         15.82           1266         TYR         CB         -5.188	1234 1235 1236 1237 1238 1239 1240 1241 1242 1243 1244 1245 1246 1247 1248 1249 1250 1251 1252	LYS LYS LYS LYS LYS LYS THR THR THR THR THR TYR TYR	CA C O CB CCD CE NZ N CA C O CB OG1 CG2 N CA C O	10.548 54.069 31.550 21.26 10.289 54.430 30.108 20.78 10.038 53.563 29.266 22.79 -9.402 54.433 32.459 22.27 -9.721 54.009 33.868 22.36 10.326 55.177 34.630 25.91 10.825 54.813 36.002 28.42 11.309 56.052 36.589 30.51 10.327 55.686 29.733 19.62 10.249 56.059 28.237 20.98 -8.822 55.510 27.732 20.84 -8.780 55.122 26.657 22.93 -9.822 57.568 28.309 19.41 -8.833 57.806 29.314 22.97 11.069 58.348 28.636 24.19 -7.719 55.518 28.646 20.25 -6.458 55.219 28.007 18.72 -5.770 53.922 28.602 18.44 -4.685 53.476 28.203 20.29
1268 TYR CD2 -5.158 54.491 32.606 21.69 1269 TYR CE1 -5.830 52.813 34.755 19.08 1270 TYR CE2 -5.580 55.003 33.823 22.03 1271 TYR CZ -5.883 54.185 34.902 20.70 1272 TYR OH -6.221 54.746 36.125 20.35 1273 THR N -6.777 50.103 31.213 13.79 1274 THR CA -7.774 49.326 31.968 14.01 1275 THR C -7.068 48.551 33.024 13.51 1276 THR O -6.050 47.935 32.736 12.55 1277 THR CB -8.369 48.336 30.922 15.24 1278 THR OG1 -9.115 48.941 29.853 15.88 1279 THR CG2 -9.278 47.291 31.615 12.04 1280 VAL N -7.636 48.525 34.247 13.29 1281 VAL CA -7.079 47.585 35.204 14.30 1282 VAL C -8.024 46.398 35.369 14.29 1283 VAL O -9.217 46.546 35.469 15.36	1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266	TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	CG CD1 CD2 CE1 CE2 CZ OH N CA C O CB	-4.89756.88829.31413.33-5.46557.86430.08114.09-3.68256.42729.72113.30-5.01658.19131.31117.45-3.09056.86630.87419.22-3.76757.71531.69619.65-3.07958.07832.80120.24-6.46153.35129.59115.06-5.91552.15130.22814.61-7.01051.38830.96215.66-8.02351.95631.32017.43-4.71952.52231.13415.82-5.11853.11532.44319.10
	1268 1269 1270 1271 1272 1273 1274 1275 1276 1277 1278 1279 1280 1281 1282	TYR TYR TYR TYR TYR THR THR THR THR THR VAL VAL VAL	CD2 CE1 CE2 CZ OH N CA C O CB OG1 CG2 N CA C	-5.158 54.491 32.606 21.69 -5.830 52.813 34.755 19.08 -5.580 55.003 33.823 22.03 -5.883 54.185 34.902 20.70 -6.221 54.746 36.125 20.35 -6.777 50.103 31.213 13.79 -7.774 49.326 31.968 14.01 -7.068 48.551 33.024 13.51 -6.050 47.935 32.736 12.55 -8.369 48.336 30.922 15.24 -9.115 48.941 29.853 15.88 -9.278 47.291 31.615 12.04 -7.636 48.525 34.247 13.29 -7.079 47.585 35.204 14.30 -8.024 46.398 35.369 14.29

1326       LEU       C       -4.851       31.471       40.119       16.97         1327       LEU       O       -4.491       31.577       41.289       13.98         1328       LEU       CB       -3.391       33.307       39.131       13.18         1329       LEU       CG       -2.359       32.718       38.152       14.58         1330       LEU       CD1       -2.685       31.494       37.300       13.64         1331       LEU       CD2       -0.889       32.995       38.428       15.94         1332       GLU       N       -5.312       30.327       39.601       18.45         1333       GLU       CA       -5.299       29.116       40.366       18.22         1334       GLU       C       -4.243       28.118       39.888       19.24         1335       GLU       O       -4.250       27.596       38.789       17.3         1336       GLU       CB       -6.667       28.458       40.354       18.1
1337 GLU CG -6.781 27.377 41.470 20.3

1338 1339	GLU GLU	CD OE1	-8.058 26.570 41.332 22.37 -8.995 26.977 40.666 21.04
1339 1340 1341 1342 1343 1344 1345 1346 1347 1348 1349 1350 1351 1352 1353 1354 1355 1356 1357 1358 1360 1361 1362 1363 1364 1365 1366 1377 1378 1379 1380 1381 1382 1383 1384 1385 1386 1387	GLN N N N N N N U U U U U U U U U R R R R	OE1 ON C C O C C O C C C C C C N C C O C O C	-8.995 26.977 40.666 21.04  -8.108 25.474 41.879 25.85 -3.345 27.828 40.804 22.37 -2.546 26.649 40.584 25.92 -3.387 25.377 40.755 25.53 -3.572 24.897 41.861 25.67 -1.325 26.765 41.500 27.66 -0.601 25.427 41.717 27.77 -1.077 24.341 41.436 27.56 0.571 25.533 42.244 29.56 -3.807 24.821 39.627 23.83 -4.652 23.629 39.641 22.45 -4.106 22.430 40.435 24.96 -4.848 21.603 40.937 26.15 -4.919 23.245 38.197 16.54 -6.200 23.824 37.584 15.77 -6.355 23.781 36.061 13.72 -6.966 24.930 38.291 12.99 -2.783 22.364 40.547 26.74 -2.184 21.257 41.276 27.87 -2.387 21.269 42.779 29.27 -2.324 20.235 43.438 30.62 -0.656 21.271 40.960 28.76 -0.222 21.276 39.580 28.70 0.076 20.197 41.798 28.43 -2.626 22.437 43.328 28.69 -2.904 22.402 44.775 28.99 -4.201 23.179 45.033 32.17 -4.673 23.210 46.146 34.29 -1.808 23.352 45.374 25.58 -1.906 24.712 44.893 24.80 -0.413 22.844 44.960 21.97 -4.726 23.838 43.975 34.19 -5.903 24.683 44.149 36.02 -5.651 25.904 45.068 34.54 -6.560 26.502 45.630 34.82 -7.056 23.797 44.633 40.16 -7.447 22.737 43.605 46.76 -8.777 22.111 44.000 50.73 -9.050 21.806 45.145 50.25 -9.586 21.893 42.985 53.71 -4.364 26.264 45.184 33.22 -4.084 27.590 45.723 33.18 -4.526 28.672 44.740 30.34 -4.574 28.507 43.524 28.93 -2.597 27.807 45.960 37.35 -2.029 27.073 47.151 45.79 -0.503 27.118 47.092 53.23 0.090 28.186 47.253 54.41
1388	GLU	OE2	0.110 26.083 46.854 58.44

1389 1390 1391 1392 1393 1394 1395 1396 1397 1398 1400 1401 1403 1404 1405 1406 1407 1418 1419 1421 1422 1423 1424 1426 1427 1428 1430 1431 1432 1433 1433	TTTTTTTTAAAAAAAAAAAGGGGGGGGLLLLLLLLLLLL	NH2 N CA C O CB CDD OE2 N CA C O CB CDD N CA	-4.771 29.824 45.341 27.29 -5.321 30.882 44.519 21.75 -4.702 32.231 44.866 20.39 -4.480 32.572 46.024 21.88 -6.901 30.855 44.831 20.91 -7.887 30.317 43.926 18.30 -7.402 32.239 45.242 22.13 -4.478 33.010 43.805 17.74 -3.917 34.344 44.023 15.94 -4.593 35.365 43.131 15.58 -4.922 35.078 41.994 14.33 -2.434 34.314 43.653 16.13 -1.673 33.304 44.487 18.13 -0.197 33.503 44.382 18.87 0.471 32.485 45.132 20.71 1.666 32.149 44.780 21.27 2.351 32.800 43.866 19.60 2.163 31.120 45.349 22.46 -4.737 36.564 43.664 16.20 -5.232 37.648 42.820 17.96 -4.066 38.466 42.226 18.25 -3.223 39.023 42.925 19.75 -6.054 38.623 43.649 20.78 -6.513 39.851 42.845 23.52 -6.777 40.984 43.840 31.18 -5.867 41.331 44.491 35.95 -7.849 41.495 43.990 33.07 -4.131 38.557 40.902 16.64 -3.169 39.310 40.109 12.04 -3.880 40.515 39.527 11.07 -4.862 40.403 38.819 11.62 -2.704 38.444 38.926 9.49 -2.292 37.027 39.359 8.19 -1.679 39.155 38.027 6.71 -1.062 36.953 40.271 8.86 -3.314 41.663 39.796 12.20 -3.765 42.894 39.153 11.34 -3.068 43.084 37.797 10.98 -1.855 43.017 37.679 11.20 -3.376 43.996 40.135 11.87 -4.459 44.428 41.136 14.45 -3.887 43.396 40.135 11.87 -4.459 44.428 41.136 14.45 -3.887 43.335 36.774 11.21 -3.409 45.000 35.405 10.38
1429	LEU	CG	-4.459 44.428 41.136 14.45
1430	LEU	CD1	-3.851 45.049 42.374 13.48
1431	LEU	CD2	-5.510 43.382 41.494 14.58
1432	HIS	N	-3.887 43.335 36.774 11.21